

# **New Insights into the Regulated Differentiation of Intercalated Cells in Mouse Kidney from Ae1 (Slc4a1) Deficient Mice**

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## **ZUSAMMENFASSUNG**

Die Niere der Säugetiere ist ein lebenswichtiges Organ, das eine Vielfalt von unterschiedlichen regulatorischen Funktionen erfüllt, wie z.B. die Aufrechterhaltung des systemischen Wasser-, Elektrolyte- und Säure-Basen Haushalts. Das renale Sammelrohrsystem setzt sich aus mindestens drei verschiedenen Zelltypen zusammen, die für die Feinabstimmung des finalen Urins verantwortlich sind. Dabei übernehmen die Hauptzellen (HZ) die Regulation von Wasser und Elektrolyten, wohingegen verschiedenen Arten von interkalierenden Schaltzellen (SZ) die Ausscheidung und Rückgewinnung von Säure-Basen Äquivalenten zukommt. Wie die Namensgebung andeutet, repräsentieren die Hauptzellen den am häufigsten vorkommenden Zelltypus im renalen Sammelrohrsystem und sind gelegentlich mit unterschiedlichen Typen von Schaltzellen (Typ A-SZ, Typ B-SZ) durchsetzt. Neben markanten Unterschieden in ihrer ultrastrukturellen Morphologie können die verschiedenen Zellen des renalen Sammelrohrs relativ einfach anhand von immunologischen Markern unterschieden werden. In der vorliegenden Studie werden Hauptzellen daher anhand eines immunologischen Nachweises des Wasserkanals Aqp2 (Aquaporin 2) identifiziert. Desweiteren werden die Säure sezernierenden Typ A-SZ von den Bikarbonat sezernierenden Typ B-SZ unterschieden, da diese beiden Typen am häufigsten im Sammelrohr vorkommen und bisher am ausführlichsten beschrieben wurden. Der immunologische Nachweis der V-ATPase (Protonenpumpe,  $\alpha 4$  und B1 Untereinheit) dient im Allgemeinen als gemeinsamer Marker für Typen von Schaltzellen. Ferner gibt die subzelluläre Lokalisierung der V-ATPase Aufschluss über den Typus der SZ. Jedoch können die beiden Haupttypen der Schaltzellen eindeutig anhand der immunologischen Marker Ae1 (Anionen Austauscher 1, Slc4a1) für Typ A-SZ und Pds (Pendrin) für Typ B-SZ identifiziert werden.

Intensive Studien der letzten zwanzig Jahre haben stetig dazu beigetragen die komplexen Prozesse induktiver und reziproker Signalwege der metanephrischen Nierenentwicklung zu entschlüsseln und zu verstehen. Auch das ausgereifte gesunde Organ wurde bisher intensiv studiert und charakterisiert. Leider ist nur sehr wenig über die Abläufe, Signalwege und genetischen Programme bekannt, die für eine regulierte Enddifferenzierung und Reifung des sich entwickelnden Organs wichtig sind. Glücklicherweise haben einige kürzlich veröffentlichte Studien an verschiedenen Modellorganismen ein wenig Licht ins Dunkel gebracht. Verschiedene Modulatoren aus den p53 und Notch Signalwegen, sowie die Transkriptionsfaktoren Foxi1 und Cp2l1 konnten letztendlich in Zusammenhang mit der terminalen Differenzierung des renalen Sammelrohrsystems gebracht werden.

Menschen und Mäuse, denen Ae1 in der Niere fehlt, entwickeln eine distale renale tubuläre Azidose (dRTA). Es ist bekannt, dass Störungen des Säure-Basen Haushalts eine adaptive Antwort der Niere induzieren, die auch eine morphologische Umwandlung des Nephron und Sammelrohrsystem zur Folge hat. In diesem Sinne führt eine chronische Säurebelastung zur Reduktion von Bikarbonat sezernierenden Typ B-SZ und gleichzeitigem Anstieg der Säure sezernierenden Typ A-SZ, wobei die individuelle und Zelltyp charakteristische Expression molekularer Marker (z.B. Ae1, Pds) unverändert bleibt. Interessanterweise konnte in der vorliegenden Studie eine ungewöhnliche Expression der Marker Pds und Aqp2 in den inneren medullären Sammelrohren von Ae1 defizienten Nieren mittels Immunfluoreszenz beobachtet werden. Hier wurde eine erhöhte Zahl an Pds exprimierenden Zellen sowie Aqp2/Pds coexprimierende Zellen im Nierenmark von Ae1 Knockout (KO) Tieren beobachtet. Dies steht im krassen Gegensatz zu Ae1 Wildtyp (WT) Nieren, bei denen eine Expression von Pds im Nierenmark nicht mehr vorkommt. Desweiteren konnten veränderte Expressionsmuster der Transkriptionsfaktoren Foxi1 und Cp2l1 in Ae1 defizienten Mausnieren festgestellt werden.



Gleichfalls wurde eine veränderte Expression von molekularen Markern der Schaltzellen in einer Nierenbiopsie eines Patienten mit der AE1 Mutation S613F beobachtet. Bei Untersuchungen zweier weiterer Mausmodelle mit unvollständiger dRTA (Knockout von Atp6v1b1 oder Rhcg) konnten derartige Veränderungen der Differenzierung von Zellen des Sammelrohres jedoch nicht festgestellt werden. Ein ähnlich negatives Ergebnis, in Bezug auf die terminale Differenzierung von Sammelrohrzellen, wurde in Nieren von Mäusen mit induzierter chronischer metabolischer Azidose über 8 Wochen gefunden. Daher kann geschlussfolgert werden, dass nicht die begleitende Azidose (dRTA) verantwortlich für die beschriebene abnorme Differenzierung des Sammelrohrs in Ae1 KO Mäusen ist, sondern das physische Fehlen von Ae1 *per se*.

Die vorliegende Studie bietet daher einige neue und interessante Erkenntnisse im Hinblick auf die terminale Differenzierung des renalen Sammelrohrsystems, die für die normale Entwicklung bzw. Regulation dieses Nephronabschnittes von Bedeutung sind. Es konnte gezeigt werden, dass die Transkriptionsfaktoren Foxi1 und Cp2l1 in unterschiedlichen Zelltypen des renalen Sammelrohrsystems exprimiert sind, was auf eine Modulation der terminalen Differenzierung durch benachbarte Zellen hindeutet. Im Einklang mit seiner Funktion als Regulator der Transkription verschiedener molekularer SZ Marker, wurde Foxi1 hauptsächlich in V-ATPase exprimierenden Schaltzellen gefunden. Cp2l1 hingegen wurde hauptsächlich in Aqp2 exprimierenden Hauptzellen detektiert, obwohl ein Fehlen von Cp2l1 auch in einem kompletten Verlust von renalen Schaltzellenmarkern (V-ATPase, Ae1, Pds) resultiert. Die Notch regulierte Signalkaskade scheint die Entwicklung der verschiedenen Zelllinien im renalen Sammelrohr der Maus zu dirigieren, vermutlich durch laterale Inhibition. Während eine Aktivierung von Notch die Präsenz von Hauptzellen in der Niere fördert, führt eine Inaktivierung von Notch zur Differenzierung von Nebenzellen.

Unter Einbeziehung der Beobachtungen an der sich entwickelnden Haut von Amphibien, der beschriebenen Resultate aus verschiedenen Mausmodellen und der

Resultate der vorliegenden Arbeit kann daher ein neues Modell für die terminale Differenzierung von Sammelrohrzellen in der Mausniere vorgeschlagen werden. Dementsprechend führt eine Aktivierung von Notch, vermutlich durch p53, zur Differenzierung von Hauptzellen aus renalen Vorläuferzellen und der Aktivierung von Cp2l1. Gleichzeitig führt die Aktivierung von Cp2l1 zur Inaktivierung von Notch in benachbarten Zellen, wodurch wiederum eine Aktivierung von Foxi1 erfolgt, welches eine Differenzierung der Schaltzellen fördert. Die Expression von Cp2l1 im Nierenmark von Ae1 KO Mäusen war gestört. Dies könnte darauf hinweisen, dass eine reziproke Interaktion oder Feedback-Schleife von Cp2l1 mit einem anderen unbekannten Schlüsselmolekül und Signalweg, oder eine negative Autoregulation von Cp2l1, im weiteren Verlauf erforderlich ist. Diesem Modell fehlt ein weiteres Schlüsselmolekül, das die Differenzierung eines bestimmten Typs von Schaltzellen interaktiv regulieren kann, da Foxi1 sowohl die Transkription von Ae1 als auch Pds reguliert. Die Ergebnisse der vorliegenden Studie unterstützen die Annahme, dass es sich bei diesem Schlüsselmolekül um den  $\text{Cl}^-/\text{HCO}_3^-$  Austauscher Ae1 selbst handeln könnte oder Ae1 damit interagiert. Zumindest scheint die Präsenz von Ae1 eine wichtige Rolle bei der Aufrechterhaltung eines bestimmten Typus der Schaltzellen zu spielen, da die Expression von Pds und Aqp2 im inneren Mark Ae1 defizienter Nieren gestört ist. Diese Hypothese wird auch dadurch gestützt, dass zuerst V-ATPase und Ae1 exprimierende Zellen in der embryonalen Niere detektiert werden und erst später Pds positive Zellen. Das bedeutet, dass sich Typ A-SZ in der zeitlichen Abfolge vor den Typ B-SZ entwickeln. Ae1 oder ein auf Ae1 basierender Multi-Protein Komplex (z.B. mit CaII) könnte also das vorangehende Modell komplettieren. Demzufolge würde eine Aktivierung von Foxi1 zur Expression von Ae1 in Typ A-SZ führen. Dies wiederum könnte eine reziproke Interaktion von Ae1 allein oder einem Ae1-Multi-Protein Komplex mit Foxi1 oder einem anderen unbekannten Molekül induzieren, um die Differenzierung von benachbarten Typ B-SZ im weiteren Verlauf zu regulieren und gleichzeitig den Typus der A-SZ in der Effektorzelle aufrechtzuerhalten.

## SUMMARY

The mammalian kidney is an essential organ that serves a variety of distinct regulatory functions, such as maintenance of systemic water, electrolyte and acid-base homeostasis. The fine tuning of the final urine takes place in the renal collecting duct (CD) system, which consists of at least 3 different types of cells. The principal cells (PC) are mostly responsible for renal water and electrolyte handling, while several types of intercalated cells (IC) regulate the excretion and recovery of acid-base equivalents. As indicated by their name, principal cells represent the major CD cell type and are occasionally interspersed with distinct types of intercalated cells (type A-IC, type B-IC). Besides their marked ultrastructural morphologic differences, cells of the renal CD are easily distinguished by immunologic markers. Hence, the present study focuses on the identification of PCs by the immunologic presence of the water channel aquaporin 2 (Aqp2). Furthermore, acid secretory type A-ICs are distinguished from bicarbonate secretory type B-ICs, since these two types of ICs represent the most abundant and best characterized IC phenotype. The immunologic presence and subcellular localization of the V-ATPase ( $\alpha 4$  and B1 subunit) is a common marker for the IC phenotype. However, the two major types of ICs are best distinguished using the immunologic markers anion exchanger 1 (Ae1, Slc4a1) for type A-ICs and pendrin (Pds) for type B-ICs.

The complex processes of inductive and reciprocal signalling required for initial metanephric kidney development were intensively studied over the past two decades steadily improving our basic understanding of nephrogenesis. Similarly, the adult and healthy organ is relatively well studied and characterized. However, the signalling events and regulatory genetic networks necessary for terminal differentiation and maturation of the developing kidney are still a black box as only very little is known. Lately, few studies with different model organisms shed some light into the dark and revealed a role for transcriptional regulators of the p53 and Notch signalling pathway,

as well as the transcription factors Foxi1 and Cp2l1 in terminal differentiation of the renal CD system.

Humans and mice deficient for renal Ae1 develop distal renal tubular acidosis (dRTA). Acid-base disturbances are known to induce adaptive responses in the kidney involving the remodelling of the nephron and collecting duct system. In addition, chronic acid challenges promote the reduction in number of base secretory type B-ICs, while the relative abundance of type A-ICs increases concomitantly. Nevertheless, the individual characteristics of the IC phenotype are preserved under these circumstances. However, in kidneys of mice with Ae1 deficiency related dRTA, an abnormal expression of Pds and Aqp2 was detected in the inner medullary collecting ducts by immunofluorescence. In contrast to wildtype (WT) mouse kidneys, Pds expressing cells and unusual Aqp2/Pds coexpressing cells were increased in abundance in the inner medulla of Ae1 knockout (KO) mice. Furthermore, the expression pattern of the transcription factors Foxi1 and Cp2l1 was altered in Ae1 deficient mouse kidneys.

Consistent with the abnormal expression of CD cell marker in Ae1 deficient inner medullary collecting ducts, we found an altered expression of molecular IC markers in a renal biopsy of a patient with the AE1 S613F mutation.

Investigations of two mouse models with incomplete dRTA (Atp6v1b1 deficiency and Rhcg deficiency) did not show any signs of abnormal CD cell differentiation. Likewise, chronic metabolic acidosis induced for eight weeks in wildtype mice did not alter the terminal differentiation of renal CD cells in mice. Hence, the accompanying dRTA or acidosis is most likely not responsible for the observed abnormal CD cell differentiation of Ae1 deficient mice, but the physical lack of Ae1 *per se*.

The present study provides some interesting and unexpected new aspects regarding the terminal differentiation of the renal CD system. Surprisingly, we localized the transcription factors Foxi1 and Cp2l1 in distinct cell types of the renal CD system, suggesting a modulation of the terminal differentiation of adjacent cells. Consistent

with its regulatory role in molecular IC markers, Foxi1 was localized in V-ATPase expressing ICs. However, Cp2l1 was mostly observed in Aqp2 expressing PCs, although Cp2l1 deficiency results in loss of characteristic molecular markers (Ae1, V-ATPase) of renal ICs. Notch signalling seems to determine the cell lineage of renal CDs in mouse, probably by lateral inhibition. While activated Notch favours the presence of PCs, inactivated Notch promotes the differentiation of renal ICs. Considering observations from amphibian skin development, other mouse models and results from the present study, we suggest the following new model of terminal CD cell differentiation in mouse kidney. Activation of Notch, probably by p53, leads to the differentiation of PCs from renal precursor and may activate Cp2l1. Active Cp2l1 might promote the inactivation of Notch in adjacent cells, which in turn activates Foxi1 and IC differentiation. Cp2l1 was abnormally expressed in the renal inner medulla of Ae1 deficient mice. Therefore, a reciprocal interaction or feedback loop of Cp2l1 with another combinatorial unknown key molecule and signalling pathway or by sequential negative autoregulation seems necessary. However, another key regulatory molecule that determines the type of ICs is missing in this model. We provide data that suggest the  $\text{Cl}^-/\text{HCO}_3^-$  exchanger Ae1 as a required key protein in the induction or maintenance of terminal differentiation of renal ICs. During embryonic kidney development, V-ATPase and Ae1 expression are detected before Pds, suggesting that A-ICs differentiate before type B-ICs. Completing the proposed model, Ae1 itself or a multiprotein-Ae1 complex (i.e. with CaII) might reciprocally interact with Foxi1 or (an)other transcriptional regulator(s). Hence, activated Foxi1 would primarily favour the differentiation of type A-ICs, which are characterized by their expression of Ae1. The multi-protein complex or Ae1 alone might then reciprocally interact with Foxi1 or another unknown molecule, which in turn regulates the differentiation of neighbouring type B-ICs and simultaneously maintains the A-IC phenotype in the effector cell.

## 1. INTRODUCTION

Multicellular organisms have the ability to maintain their systemic pH within narrow ranges, despite constant local and systemic changes of acid and base equivalents due to metabolic and catabolic processes. A stable *milieu interieur* is essential to ensure an optimal (working) environment for all chemical and bio-physical processes in the organism. In humans, the pH of most extracellular fluids ranges from 7.35 – 7.45 and is tightly regulated by two buffering systems, the volatile and non-volatile buffers. By definition, buffers are substances in aqueous solutions that are able to reversibly accept or donate hydrogen ions ( $H^+$ ) and thus stabilize a given pH. In mammals systemic pH is maintained by non-volatile buffers like calcium carbonate, phosphate, hemoglobine and other proteins, uric acid and ammonium. However, the most important buffer system is the volatile carbon dioxide/bicarbonate buffering system, which is enabled by the action of the lungs and kidneys. The constant exhalation of carbon dioxide ( $CO_2$ ) by the lungs and the renal regulation of bicarbonate ( $HCO_3^-$ ) makes this buffer pair a powerful and theoretically infinite, open buffering system.

The pH dependency on the  $pCO_2/HCO_3^-$  ratio is described by the Henderson-Hasselbalch equation (Eqn. 1). Accordingly, changes in blood pH are caused by shifts in the partial pressure of  $CO_2$  ( $pCO_2$ ) in blood, or by variations of blood bicarbonate concentrations.

$$\text{Eqn. 1} \quad pH = 6.1 + \log \frac{[HCO_3^-]}{0.03 \times pCO_2}$$

Carbon dioxide is a waste product of cell respiration and reacts slowly with water to form carbonic acid, which rapidly dissociates to a proton and bicarbonate (Eqn. 2). The hydration reaction is usually accelerated and catalyzed by isoforms of the enzyme carbonic anhydrase.



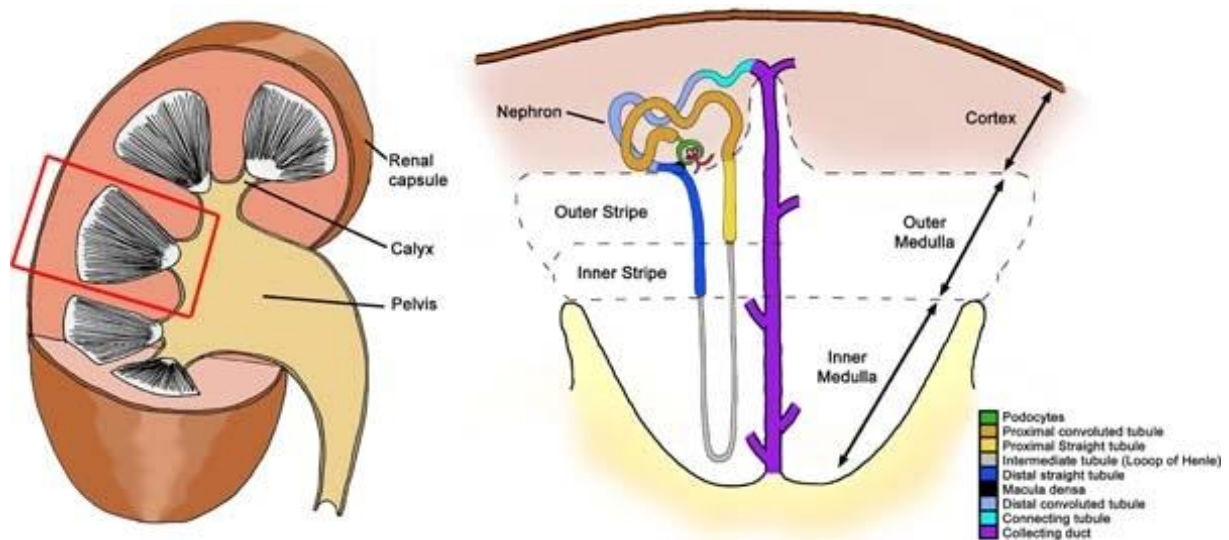
In mammals volatile  $\text{CO}_2$  is a huge source of daily acid production (15,000 mmol  $\text{CO}_2$ /day) [1]. However, since the lungs continuously remove this potential acid by regulated exhalation, this amount is less relevant in healthy individuals. Importantly, dietary metabolism and daily loss of base equivalents into stool lead to the accumulation of non-volatile acids (70 mmol/day) [1], which essentially consume plasma bicarbonate and thus need to be removed from the body. Here, the kidneys play the essential role in the removal of this daily load of non-volatile acids by recovering virtually all filtered bicarbonate as well as generating new bicarbonate. The kidneys generate new bicarbonate in two ways: the secretion of protons along with titratable acids (i.e. phosphate, citrate) and the synthesis and excretion of ammonium.

## **2. MAMMALIAN KIDNEY STRUCTURE AND DEVELOPMENT**

### ***2.1 Organization of the mature mammalian Kidney***

The mammalian kidney is a paired organ fulfilling a variety of essential functions. Besides the removal and excretion of metabolic waste products from the blood, it plays a central role in systemic acid-base balance, as well as maintaining salt and water homeostasis thereby regulating systemic blood pressure. The functional unit of the kidney is the nephron, which is organized in different segments (Fig. 2.1). In humans, each kidney consists of 800.000 – 1.200.000 nephrons, whereas in mouse approximately 11.000 nephrons are present [1, 2]. Each nephron begins with a glomerulum, which is a network of blood capillaries and podocytes that filter blood. The glomerulum is surrounded by the Bowman capsule that extends into an epithelial tubular structure (renal tubule) and finally connects to the collecting duct system (CDS). The nephron is subdivided into three domains, the proximal tubule (PT), the loop of Henle (LOH) and the distal tubule (DT). Based on ultrastructural and functional properties these domains are further distinguished into the early PT, the proximal convoluted and straight tubule, thin descending and ascending limbs of LOH, thick ascending limb of LOH, as well as distal convoluted tubule (DCT) and connecting tubule (CNT). Likewise, the renal collecting duct system shows a segmental organization and cortical collecting duct (CCD), outer medullary collecting duct (OMCD) and inner medullary collecting duct (IMCD) are distinguished.





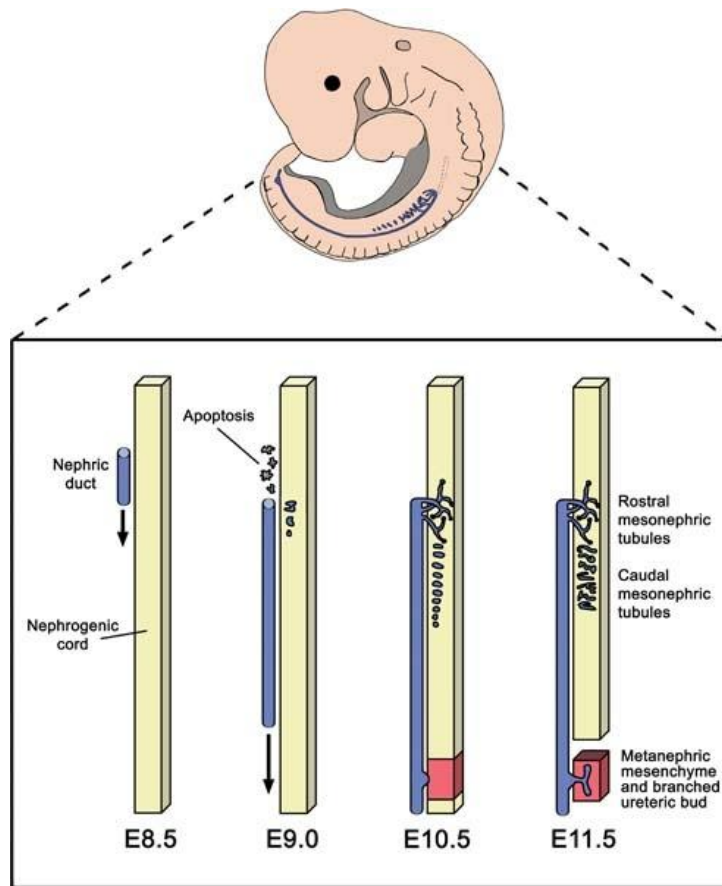
**Fig. 2.1 Structure of the mammalian kidney**, taken from [2]. Each kidney is surrounded by a fibrous renal capsule. The kidney is organized into the peripheral renal cortex and the renal medulla, which is arranged in multiple pyramidal structures. The tip of each pyramid (papilla) ends into minor and major calyces that empty into the renal pelvis. The renal pelvis is connected to the ureter which transmits the urine into the bladder. Image enlargement shows segmental organization of the nephron and collecting duct [2].

## 2.2 Development of the mammalian Kidney

The development of the mammalian kidney (metanephros) is unique, as it originates from two distinct embryonic precursor populations, the metanephric mesenchymal blastema and the ureteric bud epithelia [3]. Hence, morphogenetic and functional development of the kidney requires complex inductive interactions of the distinct precursor populations and their differentiation pathways are tightly regulated in time and space by different gene products. Among them are transcription factors, protooncogenes, growth factors, signalling molecules, and receptors. However, kidney morphogenesis is also dependent on cell adhesion molecule complexes (CAM) and their associations with the cytoskeleton, extracellular matrix (ECM) glycoproteins and ECM receptors like the integrins, as well as ECM degrading proteases [4]. Mammalian kidney development is further characterized by the formation of a more complex organ in three successive phases: the rudimentary pronephros, the transitory

mesonephros, and the permanent metanephros. In mammals and birds, the pronephros and mesonephros are transient structures, which degenerate or become modified to serve as a part of the male genital tract [5], while the metanephros differentiates into the permanent kidney. However, the pronephros is the functional filtering unit in larvae of amphibians and fish, while the mesonephros is the functional kidney in adults [6]. Thus, the three kidney forms of amniotes reflect the evolutionary development of the kidney approved by conserved genetic programming among different species [7].

The metanephros is the most complex structure and displays the functional kidney of adult mammals and birds. The development of the mammalian metanephric kidney is initiated at week 4-5 in humans and embryonic day 8.5 (E8.5) in mice [2, 4]. It begins with the formation of the pronephric duct (Wolffian duct) and mesonephric tubules from the intermediate mesoderm (Fig. 2.2). As they fuse and migrate into the adjacent metanephrogenic mesenchyme (MM), the ureteric bud (UB) emerges at the tip.



**Fig. 2.2 Initiation of kidney development in mouse**, taken from

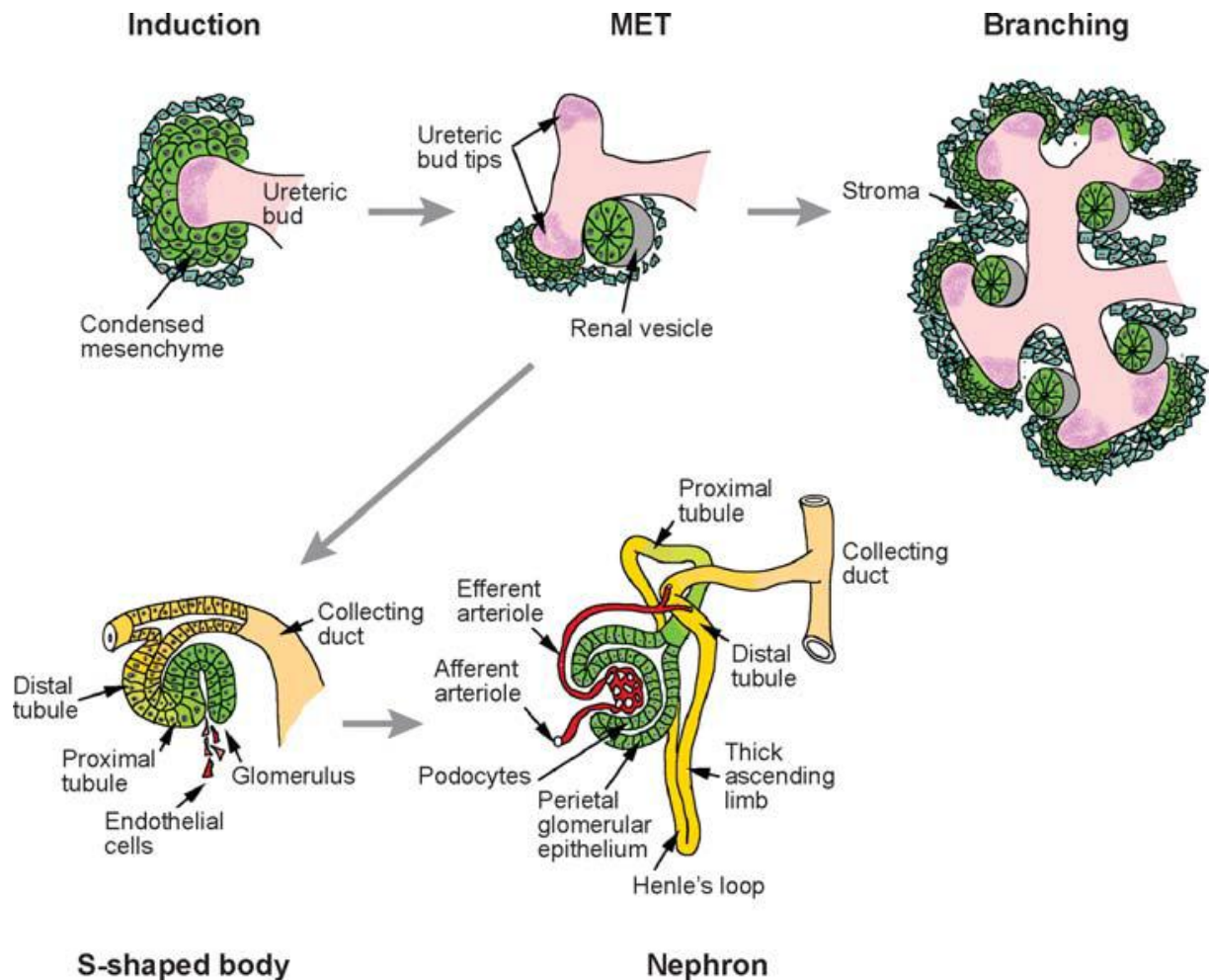
[2]. In mouse, kidney development starts at E8.5 with the formation of the nephric (Wolffian) duct primordium (blue) from the nephrogenic cord (intermediate mesoderm, yellow). Caudal nephric duct precursors grow towards the cloaca, while rostral cells become apoptotic. The extending nephric duct induces the formation of mesonephric tubules in the adjacent mesoderm. Rostral mesonephric tubules fuse to the nephric duct, while the ureteric bud (UB) emerges from the nephric duct near the cloaca at E10.5. The UB invades into a

specialized population of nephrogenic cord cells (red), the metanephric mesenchyme (MM). Reciprocal inductive interactions between UB and MM lead to repeated branching of UB and the formation of metanephric nephrons from MM (not shown).

Reciprocal inductive interactions between the UB and MM are the key events of mammalian nephrogenesis (Fig. 2.3) and accompanied by the sequential activation of gene expression from several different gene families [6]. The MM induces a repetitive branching morphogenesis of the UB, which gives rise to the collecting system. Simultaneously, the invading UB induces a mesenchymal to epithelial transition (MET) of the MM, forming metanephric nephrons in a stereotypic sequence of morphological stages (Fig. 2.3) [8].

### **2.3    *Formation of the Nephron***

The epithelial segments of the nephron, which comprises the glomerulum, proximal tubule, loop of Henle and the distal tubule (distal convoluted tubule and connecting tubule) arise from the metanephric mesenchymal blastema. Nephron development requires a series of events which is defined by morphological stages and can be divided into different steps: 1) condensation, 2) epithelialisation, 3) early morphogenesis and 4) tubule maturation [9]. Upon ureteric bud induction, cells of the MM start to condensate around the advancing UB, forming a 4-5-cell thick layer, the cap condensate (Fig. 2.3). A cluster of cap condensate cells near the UB ampulla and the adjacent branch starts to polarize as they undergo mesenchymal to epithelial transformation. These cells separate from the cap condensate and reorganize in a pretubular aggregate to form a renal vesicle. Pluripotent precursor cells within the renal vesicle give rise to all type of glomerular and tubular epithelial cells. Nephron segmentation into globular and tubular domains is induced by the formation of two clefts within the renal vesicle. Firstly, a comma-shaped body forms with the invagination of a lower cleft (vascular cleft). Thereafter, a second upper cleft builds up the S-shaped body, which is characterized by three segments or limbs. The upper limb gives rise to the LOH and distal tubule, the middle limb to the proximal convoluted tubule and the lower limb to the glomerulum. Angioblasts invade the vascular cleft of the S-shaped body and contribute to the glomerular capillary. Finally, the distal part (upper limb) of the S-shaped body fuses to the adjacent collecting duct, completing the formation of the nephron (detailed review by [10]).



**Fig. 2.3 Sequential steps of mammalian kidney development**, taken from [7, 10]. Induction of the metanephric mesenchyme by the ureteric bud promotes aggregation of mesenchymal cells to form a cap condensate around its tip. A cluster of cells separates from the cap condensate and forms a renal vesicle while they polarize and undergo mesenchymal to epithelial transformation (MET). Meanwhile, unpolarized stromal mesenchyme cells migrate into the interior of the developing organ. Thereby, they influence branching morphogenesis of the ureteric bud which continues to induce nephrogenesis. Individual renal vesicles fuse to the proliferating collecting ducts. By the S-shaped body stage, the nephron is already patterned along the proximal-distal axis. Invasion of the proximal cleft by endothelial cells starts the process of glomerulogenesis [7, 10].

## **2.4    *Formation of the renal Collecting System***

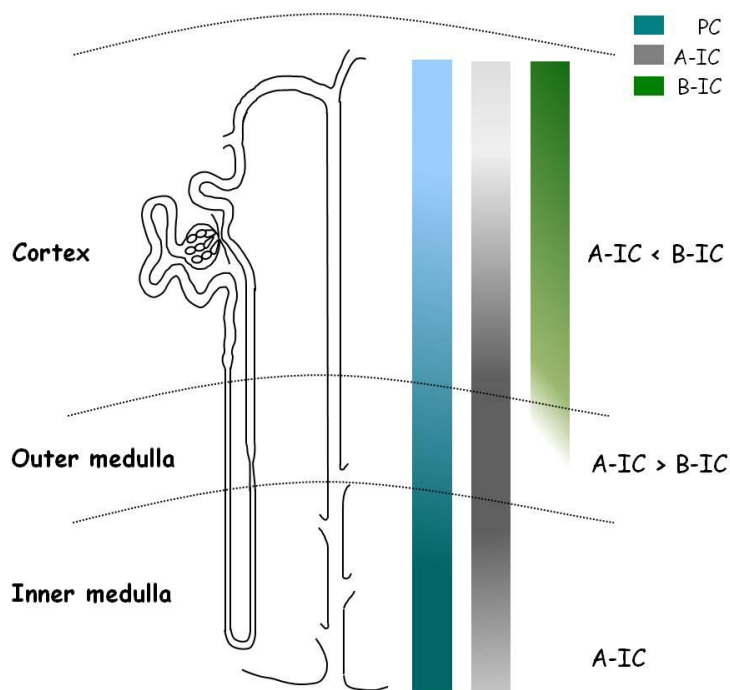
The renal collecting system comprises the cortical and medullary collecting ducts, the renal calyces, the renal pelvis and the ureter. It develops from the ureteric bud by a process called branching morphogenesis while it invades the MM. Repetitive dichotomous branching events result in several generations (~15) of UB branches, which ultimately differentiate into distinct parts of the renal collecting system. The number of UB branches elaborated is considered to be a major determinant of the final number of nephrons, since the UB branch tips (ampullae) reciprocally induce discrete subsets of MM cells to undergo nephrogenesis [10]. Thus, UB branching morphogenesis might also determine the later architecture of the mature kidney [11]. Ureteric branching is completed by weeks 20-22 of gestation in humans and at E15.5 in mouse [7, 10]. Thereafter, the collecting duct system undergoes morphologic changes that result in the patterning of cortical (peripheral) and medullary (central) domains. At this stage, distinct morphologic features emerge between the cortical and medullary collecting ducts (CDs) [12]. While cortical CDs continue to undergo branching and induction of MM, medullary CDs organize into elongated, relatively unbranched linear arrays [12]. The main trunk of the UB gives rise to the mature ureter, while the early branches undergo remodelling by increased growth and dilatation to form the renal calyces and pelvis [12, 13].

## **2.5 Terminal Differentiation and Organization of the renal Collecting Duct System**

Intensive *in vivo* and *in vitro* studies revealed many factors and molecules involved in UB outgrowth, branching morphogenesis and induction of mesenchymal to epithelial transition [14]. However, regulatory mechanisms of terminal CD system differentiation and maturation are still largely unknown. During organ development the function of the embryonic collecting duct epithelium needs to transdifferentiate from an inductive to an excretory system [15]. The architecture of the mature renal CD system is unique as it is composed of distinct cell types with highly specialized functions (see Chapter 2.6 and 3.4). Hence, the process of terminal differentiation goes along with the acquisition of functional differentiation and requires cell cycle arrest [16]. Our knowledge of the genetic programs and signals that regulate functional differentiation of the kidney has only just begun. A small number of key regulators and pathways such as the p53 and Notch signalling cascade were proposed to determine segmental nephron identity and terminal epithelial differentiation fate. Within these pathways, some transcription factors and signalling molecules (i.e. Bdkrb2, Cp2l1, Dmbt1 (Hensin), Foxi1 and Klf4) are suggested critical for the regulated differentiation of individual CD cell types. However, their hierarchical positioning within the signalling network remains to be determined. The majority of these molecules was incidentally discovered in disease models or deduced from *in vitro* studies and will be described in more detail in chapters 2.7 and 5.2.1.

The morphology of the mature renal collecting duct system in mammals is characterized by its functional, zonal and cellular heterogeneity, accounting for the functional demands of final urine processing [15]. At least three types of cells are identified based on ultrastructural features (i.e. distribution of intracellular organelles, organization of plasma membrane), molecular composition and function. Even though they are unequally distributed throughout the CD system, principal cells (PCs)

are the major cell type and responsible for water and electrolyte transport. Occasionally, the CDs are interspersed with two types of intercalated cells, type-A and type-B (A-IC, B-IC), which are involved in acid-base excretion and show a zonal distribution. In the mature mouse kidney, A-ICs are most frequently present within the outer medulla, while B-ICs are restricted to the renal cortex (Fig. 2.4) [17]. Furthermore, both types of intercalated cells are present in the late DCT and CNT [18-21]. However, based on immunologic and morphologic criteria species specific intermediate forms of intercalated cells have been described (detailed description in chapter 2.6) [22-24]. In any case, it remains controversial, whether the different ICs are truly distinct cell types or represent several configurations of the same cell [18].



**Fig. 2.4 Zonal distribution of cell populations in mouse renal collecting duct system, identified by immunologic criteria.** Within the mature renal CD system, principal cells (PCs, blue bar, left) are the major cell type. However their abundance increases from the cortex towards the inner medulla. Type A-ICs (grey bar, centre) are predominantly expressed within the outer medullary CDs, whereas type B-ICs (green bar, right) are exclusively found in the renal cortex.

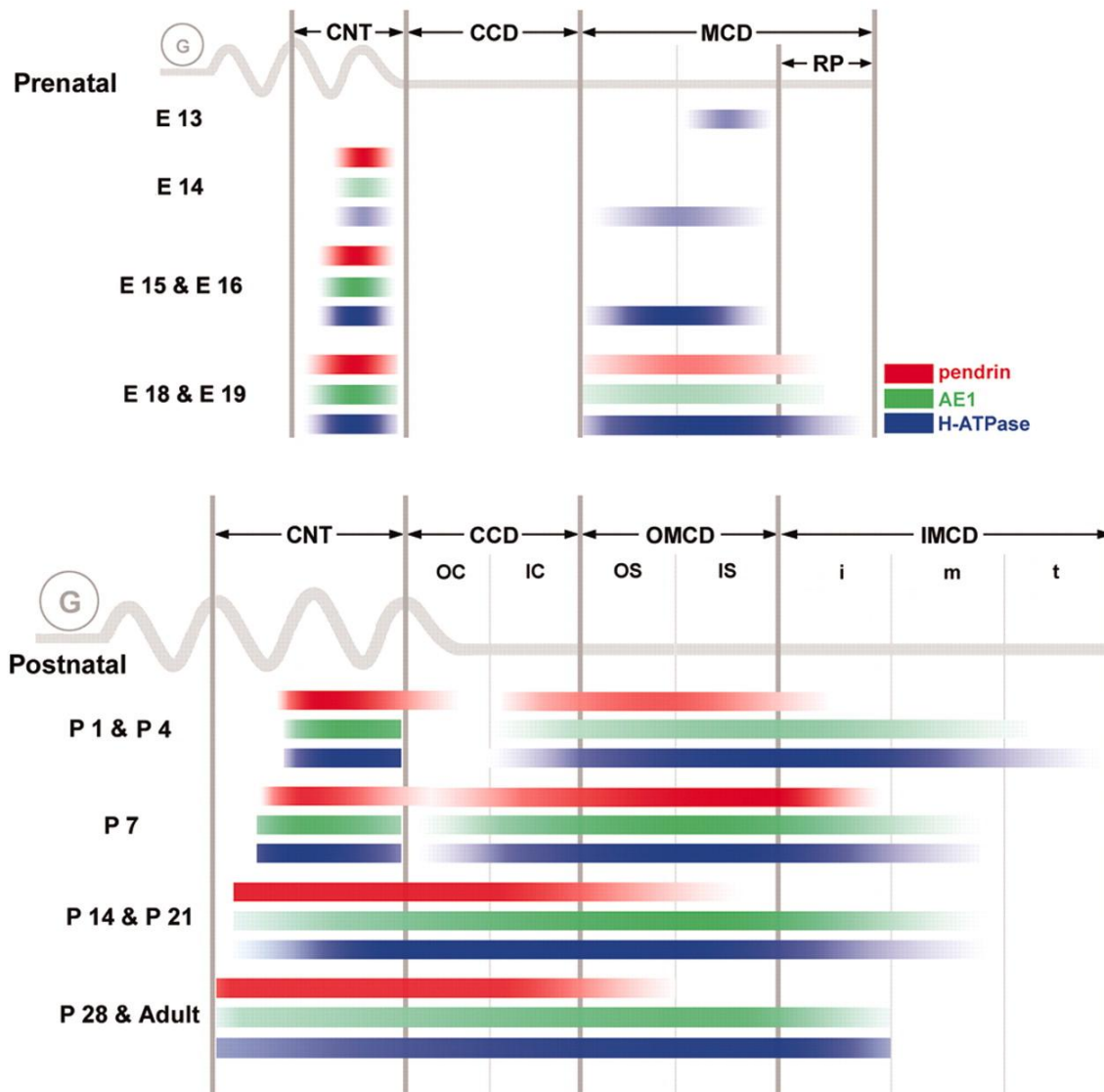


However, good evidence for the simultaneous differentiation of ICs from undifferentiated precursor populations in separate foci of the kidney in rodents was found [18, 25-27]. In mice, immunohistochemical identification of ICs demonstrated their contemporary appearance in the fetal connecting tubule and medullary collecting duct (MCD), while cortical collecting ducts appeared still not differentiated (Fig. 2.5, top). Hence, ICs seem to develop from distinct precursor populations which are present in both, the metanephrogenic blastema and the ureteric bud. Nevertheless, it remains speculative, whether PCs and the distinct types of ICs derive from the same or from distinct precursor populations.

In fetal rat kidneys immunologic marker identifying ICs are not detected before embryonic day 18 (E18) [18, 28]. In mice, renal IC specific proteins are detected the earliest at E13 and E14, respectively (Fig. 2.5) [25]. Consistently, cellular expression levels of transporters and other proteins adapt to the increasing functional demands. A sequential and segmental regulation of intercalated cell subtypes and differential expression of characteristic proteins (i.e. V-ATPase, CaII) were observed during tubule maturation (Fig. 2.5) [29, 30]. Soon after birth, the maturing renal tubule develops excretory and reabsorptive functionality and progressively achieves the morphology of the adult kidney by adaptive remodelling [31]. Hereby, ICs are gradually removed from the papillary surface epithelium and terminal MCDs, while their abundance increases erratically in the renal cortex and outer medulla, respectively (Fig. 2.5, bottom) [32]. Moreover, pendrin expressing B-ICs disappear completely from the entire mature MCDs and are restricted exclusively to the late DCT (DCT2), CNT, CCD, and early OMCD [21, 25-27, 31, 33]. Elimination of ICs from the developing rat MCD might involve two mechanisms. Type A-ICs seem to be extruded from the renal epithelium into the lumen, while type B-ICs become apoptotic and are phagocytosed by adjacent cells [32].

In human fetal kidney immature or undifferentiated ICs are observed as early as 8 weeks gestation, while characteristic AE1 expression is not detected before 14 weeks gestation. Moreover, Pendrin expression is observed only as late as 26 weeks gestation and solely restricted to the fetal renal cortex. Additionally and likewise in contrast to rodents, IC abundance in human fetal kidney increases progressively from the inner cortex to the inner medulla with type A-ICs being the predominant IC manifestation [27].

In outlook of the mature collecting duct epithelium, a structural and functional plasticity of the relative IC distribution has been observed as well in adaptive responses to disturbances of the *milieu interieur*. These observations raised the concept on reversal of structural and functional polarity in intercalated cells, with a definite proof still pending (detailed description in chapter 5.2.1) [17].



**Fig. 2.5 Segmental distribution of acid-base transporters expressed in renal intercalated cells of the developing mouse**, taken from [25]. E= embryonic kidney; P= postnatal kidney; RP= renal papilla; OC= outer cortex; IC= inner cortex; OS= outer stripe of outer medulla; IS= inner stripe of outer medulla; i= initial part of IMCD; t= terminal part of IMCD. It is of note, that  $H^+$ -ATPase and pendrin expressing cells are already detected in the late distal convoluted tubule (DCT2) of mouse, rat and human kidney (not shown) [21].

## ***2.6 Architecture and Characteristics of renal Collecting Duct Cells***

Intensive studies on the heterogeneous cytology of mammalian collecting ducts revealed numerous ultrastructural and molecular differences among cell types accounting for their distinct function. However, transitional changes of the structural appearances of each cell type are described all along the CD system, while molecular profiles and functional aspects remain rather constant. For the sake of simplicity, the present study focuses on the identification of principal cells and intercalated cells (type A and type B) by immunohistochemical markers.

### ***2.6.1 Principal Cells***

By light-microscopy, principal cells appear translucent while intercalated cells appear dark due to their high density of mitochondria. In higher magnification, the apical membrane of PCs consists of a prominent single cilium with few and short microvilli and microfoldings. The basal membrane forms infoldings (basal labyrinth), which appear as a characteristic basal rim, as it is devoid of any cell organelles. However, principal cells undergo gradual, from the deep cortex towards the upper third of the inner medulla, but species specific distinct morphological changes regarding the abundance of mitochondria, lysosomal organelles and basal infoldings. Subsequently, the cell size may increase drastically (rabbit, guinea pig) and the cell shape might change from polygonal to cuboidal or low columnar (rat, mouse) [17]. However, immunologically all variations of principal cells are identified by the luminal and sometimes basolateral expression of aquaporin 2 (Aqp2), (Fig. 2.6) [34, 35].

### **2.6.2 Intercalated Cells**

As mentioned earlier, the renal CNTs and CDs are interspersed with distinct manifestations of intercalated cells. Among the diverse IC cell population, species specific differences regarding structural manifestations, protein expression profiles and cellular pattern, as well as axial changes in their zonal distribution along the CNT and CD system have been described [26, 36-39]. All these subtypes share some common features and are involved in potassium and chloride reabsorption and acid-base homeostasis [40]. The apical membrane of ICs is usually more rounded, compared to the polygonal architecture of adjacent PCs. Intercalated cells appear darker by light microscopy because of their denser cytoplasm, which contains numerous mitochondria, polysomes and tubulovesicular membrane structures. Moreover, the luminal membrane carries elaborate microfolds or microprojections and lacks the single cilium [17]. Immunologically, intercalated cells are characterized by their expression of cytosolic carbonic anhydrase II (CaII) and a vacuolar type H<sup>+</sup>-ATPase (V-ATPase) (Fig. 2.6) [41, 42]. However, two IC manifestations are predominant and distinguished as type A and type B.

#### **2.6.2.1 Type A-Intercalated cells**

The apical cell pole of type A-ICs is usually broad and covered with elaborate microfolds and characteristic tubulovesicular structures. The apical and vesicular membrane domain is densely covered with studs. Type A-ICs are equipped with numerous mitochondria, which often arrange along the apical membrane and the nucleus shifted to the basolateral side [42]. Many microtubules and clathrin-coated vesicles are located between the tubulovesicular profile and mitochondria [17, 43]. Correspondingly, high abundance of the V-ATPase in the apical pole and membrane was demonstrated and is usually evident of type A-ICs [22, 24, 44]. The basolateral membrane of A-ICs is uncoated and devoid of V-ATPase. A kidney specific isoform of the Cl<sup>-</sup>/HCO<sub>3</sub><sup>-</sup> exchanger Ae1 (anion exchanger 1, band3, Slc4a1) is exclusively

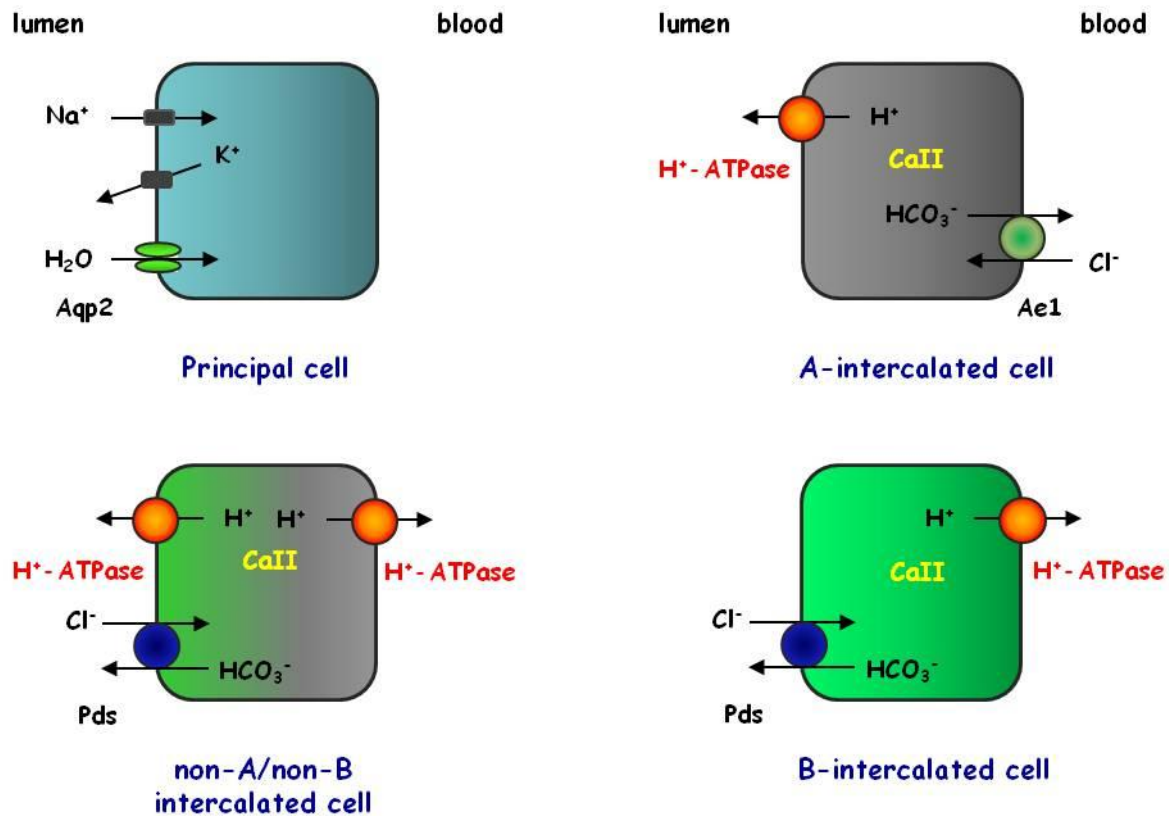
expressed in the basolateral membrane of type A-ICs and serves as an immunologic marker (Fig. 2.6) [43, 45]. Thus, A-ICs are considered as acid secretory cells and show marked hypertrophy and hyperplasia under conditions of metabolic and respiratory acidosis and potassium depletion [17].

#### **2.6.2.2      *Type B-Intercalated cells***

In many respects type B-ICs are structurally and functionally opposite to type A-ICs (as summarized in detail by [17]). Compared with type A-ICs, the apical cell pole of B-ICs is usually narrow, with few and short microvilli, while the basal pole is broad with tightly arranged deep infoldings of the membrane. B-ICs exhibit a characteristic homogeneous belt of dense vesicle-free cytoplasm, containing filamentous compounds, just below the apical membrane [42]. Thus, mitochondria usually align along the lateral membrane and vesicles are distributed throughout the remaining cytoplasm. Type B-ICs contain a large Golgi-Complex with abundant smooth endoplasmic reticulum, exhibiting clathrin-coated heads. Polyribosomes, microtubules, lysosomes and phagolysosomes are much more frequently present in B-ICs than A-ICs [17]. In contrast to type A-ICs, B-ICs usually lack apical V-ATPase expression and correspondingly apical vesicles and studs. Instead, V-ATPase and coated studs are abundant in the basolateral membrane domain [17, 42]. Lectin-immunoreactivity of the luminal membrane to peanut agglutinin (PNA) identifies B-ICs. However, a member of the Slc26 family, pendrin (Pds, Slc26a4) was finally identified as the long missing exclusive  $\text{Cl}^-/\text{HCO}_3^-$  exchanger on the luminal membrane of B-ICs [46, 47]. Thus, B-ICs serve as bicarbonate secreting cells, mirroring the action of A-ICs, allowing the fine tuning of urinary acid-base excretion [33].

### **2.6.2.3      *Intermediate forms of Intercalated Cells***

Based on morphologic criteria, intermediate variants of ICs, which cannot be attributed to type A-ICs nor type B-ICs, have been observed [17]. One manifestation of these intermediates is frequently observed in rat and mouse renal CNT and CCD and is referred to as non-A/non-B intercalated cells (non-A/non-B-IC), as they display features of both, type A-ICs and B-ICs. The non-A/non-B cells are characterized by the luminal presence of both, V-ATPase and Pds, as well as the absence of basolateral Ae1 (Fig. 2.6) [42, 46, 48]. Functionally, the equilateral expression of V-ATPase and Pds would result in the parallel secretion of  $H^+$  and  $HCO_3^-$  in the exchange for luminal  $Cl^-$ . However, it remains controversial, whether non-A/non-B cells represent a distinct cell type or a specific activity state of B-ICs [33].



**Fig. 2.6 Schematic representation of molecular collecting duct marker.**

Immunohistochemically, principal cells along the CNT and CD are identified by the luminal presence of aquaporin 2 (Aqp2). Type A-intercalated cells are characterized by the luminal expression of V-ATPase ( $\text{H}^+$ -ATPase) and basolateral expression of anion exchanger 1 (Ae1). In type B-intercalated cells the V-ATPase is expressed on the basolateral membrane, while pendrin (Pds) is found on the apical side. Type non-A/non-B intercalated cells share features of both, type A-IC and type B-IC, with luminal coexpression of V-ATPase and Pds in the absence of basolateral Ae1. Moreover, some type non-A/non-B-ICs express the V-ATPase on the apical and basolateral membrane simultaneously. All types of intercalated cells express high levels of cytosolic carbonic anhydrase II (CaII).



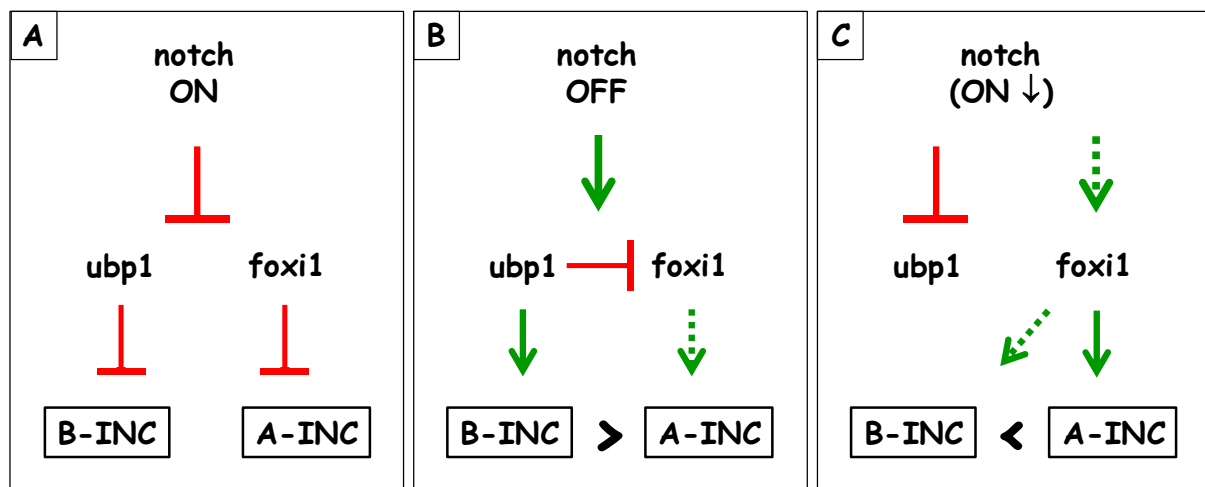
## **2.7 Disturbances of terminal renal Collecting Duct Differentiation**

Even though the pathways and modulators of late embryonic renal CD differentiation are largely unknown, a variety of distinct molecules were discovered and suggested to play a role in mouse CD differentiation and adaptive plasticity of the mature nephron. Among them are receptors, signalling molecules, transcription factors, enzymes and transport proteins [49]. Strong evidence for most of the regulatory molecules and pathways comes from mouse disease models and *in vitro* experiments. However, developmental studies of other model organisms, such as zebrafish (pronephros and epidermis) and *Xenopus laevis* (larval skin), are of great support, as they share structure-function similarities and conserved regulatory mechanisms with the mammalian renal CD system (Fig. 2.7) [50-53].

The differentiation of ionocytes (proton-secreting cells, PSC) in the larval epidermis of *Xenopus laevis* is probably the best available model, resembling mammalian renal intercalated cells. It has been shown recently, that two types of ionocytes are present and distributed in a “salt and pepper” fashion in the *Xenopus* larval skin. Similar to mammalian renal ICs, *Xenopus* type alpha-ionocytes (A-INCs) are characterized by the basolateral expression of ae1 and apical v-ATPase, while type beta-ionocytes (B-INCs) express apical pendrin and basolateral v-ATPase [51]. Moreover, differentiation of the ionocyte subtypes is regulated by notch activity and the downstream targets upstream binding protein 1 (ubp1) and foxi1 (Fig. 2.7) [51].

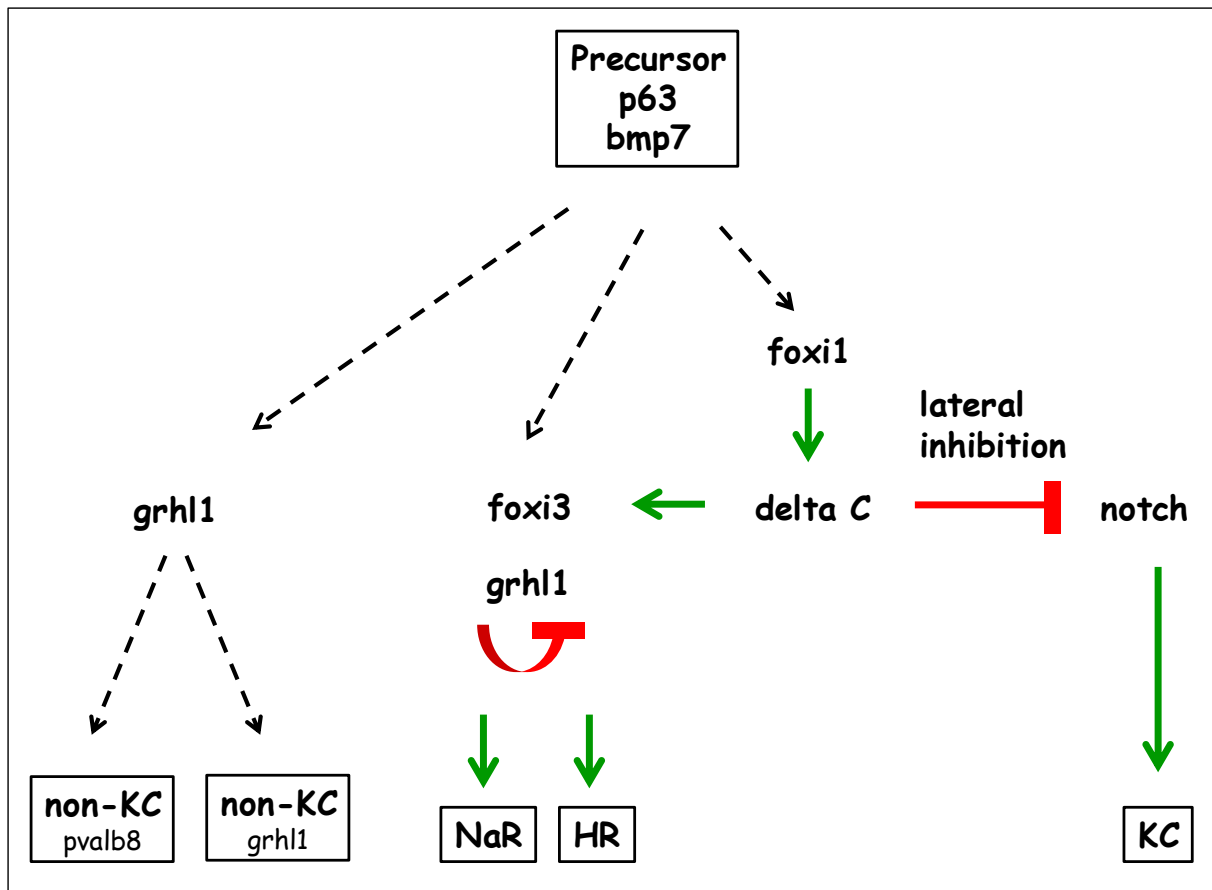
*Xenopus* ubp1 belongs to the family of grainyhead transcription factors (chapter 2.7.4) and shares close sequence similarities with the leader binding protein 1a (Lbp1a) of mouse and men. However, direct orthologues of ubp1 in mouse and men were not reported [51]. Additionally, based on its proposed functional role, *Xenopus* ubp1 rather resembles an orthologue of mammalian Cp2l1 (chapter 2.7.4) [51].

However, in *Xenopus* larval skin, expression of foxi1 was found in both types of ionocytes, whereas ubp1 seems to localize specifically with pendrin expressing B-IONCs. Interestingly, notch activity might mediate the regulation of ubp1 and foxi1 in a dose dependent manner. Strikingly, the notch signalling pathway, as well as the transcription factors Foxi1 and Cp2l1 (mammalian orthologue of ubp1) are suggested critical determinants of mammalian renal IC differentiation.



**Fig. 2.7 Suggested model of *Xenopus laevis* epidermal ionocyte differentiation.** The schematic model is based on the findings of [51]. **A)** Overexpression of activated notch leads to suppression of ionocyte (INC) formation, probably by inactivating ubp1 and foxi1. **B)** Inhibition of notch leads to the formation of ionocytes, presumably by activation of ubp1 and foxi1. However, B-IONCs (type beta-ionocytes) are favoured over A-IONCs (type alpha-ionocytes), most likely by ubp1 mediated repression of foxi1. **C)** Dose dependent effect of notch. While inactivated notch abolishes INC formation (**A**), little amounts of activated notch lead to the formation of INCs, favouring A-IONCs over B-IONCs.

Similar observations were made regarding the differentiation of distinct ionocyte populations and keratinocytes in the epidermis of zebrafish embryos and larvae [52]. Cell lineage tracing experiments revealed the development of ionocytes and keratinocytes from the same precursor cell pool [54]. At least two types of ionocytes are distinguished,  $\text{Na}^+, \text{K}^+$ -ATPase-rich (NaR) cells and  $\text{H}^+$ -ATPase-rich (HR) cells. Apparently, members of the fox transcription factor family (foxi1 and foxi3) are essential in ionocyte specification [54-56]. Furthermore, the delta-notch signalling is required to inhibit ionocyte formation of neighbouring cells, thus inducing the keratinocyte phenotype by lateral inhibition. Recently, the grhl1 (grainyhead-like 1) transcription factor was discovered to play some role in ionocyte and non-keratinocyte formation, probably by negative autoregulation (Fig. 2.8) [57].



**Fig. 2.8 Predicted model of epidermal cell differentiation in zebrafish larvae**, modified after [56, 57]. A common pool of epidermal precursor cells (expressing p63, bmp7) give rise to distinct cell types. Hereby, foxi1 mediated activation of delta C seems to determine the cell fate of neighbouring cells. While delta C mediated induction of foxi3 gives rise to the ionocyte lineage ( $\text{Na}^+$ ,  $\text{K}^+$ -ATPase-rich cells, NaR;  $\text{H}^+$ -ATPase-rich cells, HR), the activation of the delta/notch signalling pathway promotes the development of keratinocyte (KC) precursor cells by lateral inhibition. Furthermore, the induction of foxi3 and the negative autoregulation of grhl1 within the ionocyte precursor might result in the terminal differentiation of NaR and HR. In addition, the activation of grhl1 alone may lead to the development of distinct non-keratinocytes (non-KC).

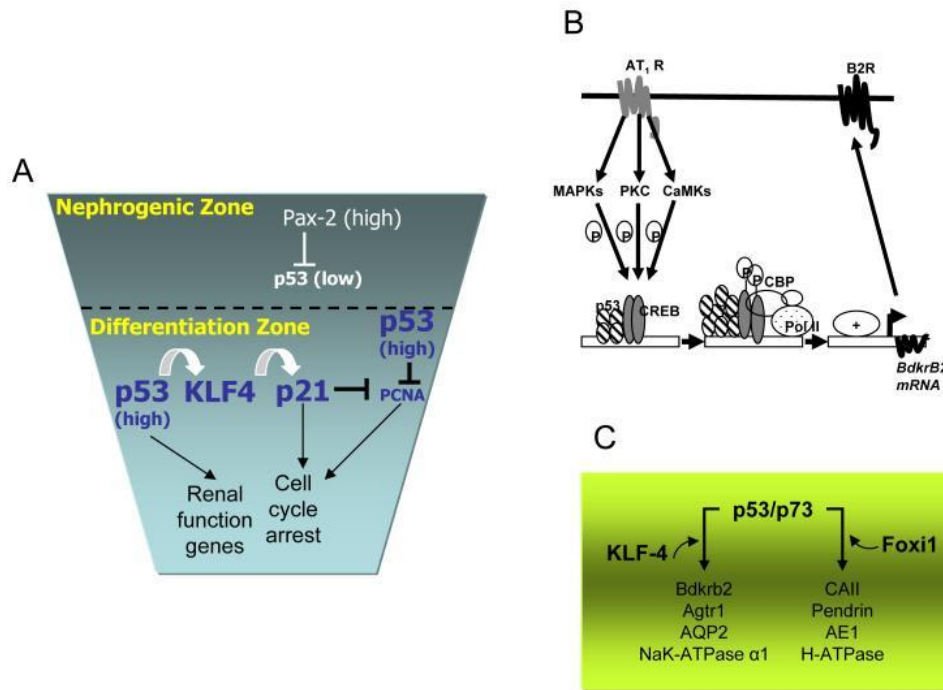
### **2.7.1 The p53 gene family**

Terminal cell differentiation inversely correlates with proliferation and thus requires cell cycle arrest and activation of a functional gene expression in the cell [16]. The tumor suppressor protein TP53 (p53) is a master regulator of transcription and central to the activation of cell cycle arrest, senescence, apoptosis, tumorigenesis, embryo development and differentiation [58, 59]. Consistently, p53 expression and activity is crucially regulated during embryonic kidney development and plays a role in transcriptional regulation of renal functional genes (RFGs) [60-62]. Briefly, in the fetal rat kidney p53 expression was detected in the differentiated zone with the onset of tubular differentiation, while it was absent in the immature nephrogenic zone [61]. The spatiotemporal distribution pattern and posttranslational modifications of p53 correlate with the expression of RFGs (i.e. Bdkrb2, Aqp2, Na<sup>+</sup>/K<sup>+</sup>-ATPase  $\alpha$ 1, AT<sub>1A</sub>) and thus differentiated renal epithelial cells [63]. Consistently, p53 was shown to control the expression of these RFGs in distinct *in vivo* and *in vitro* models [61, 63, 64]. In addition, transcription factors (i.e. Klf4, Foxi1) involved in epithelial differentiation might be controlled directly by p53 or signalling molecules within the p53 pathway [65-67]. Interestingly, p53 deficient newborn mice exhibit severe renal cysts and hyperplasia, while p53 overexpression results in renal hypoplasia [61]. Among the p53 responsive genes the bradykinin receptor type B2 (Bdkrb2, B2R) was identified and their relationship was investigated in more detail. The bradykinin receptors are thought to mediate paracrine actions of the renal kallikrein-kinin-system (KKS), which comprises cellular growth, inflammatory response, modulation of renal blood flow and urinary sodium excretion [68, 69]. Indeed, salt stressed Bdkrb2 deficient mouse embryos develop renal CD dysplasia and die early after birth [70]. The observed renal dysgenesis might be a result of p53 mediated apoptosis of nephron progenitors and repression of terminal differentiation [71, 72]. Moreover, a combinatorial model involving the transcription factors p53, cAMP response-element binding protein (CREB), Krüppel-like factor 4 (Klf4), and the transcriptional coactivator CREB-binding protein (CBP) is proposed to cooperatively activate Bdkrb2 transcription in the

process of terminal nephron differentiation (Fig. 2.9) [13, 65, 66]. Accordingly, p53 performs a dual role in terminal epithelial differentiation by controlling cell cycle arrest and activating the transcription of renal function genes. However, whether a multi-protein complex is also required for the transcriptional regulation of other p53 targets such as Aqp2 or Na<sup>+</sup>/K<sup>+</sup>-ATPase is still unknown.

In addition, p73, another member of the p53 family and close homologue, was identified to control the expression of Aqp2 and Bdkrb2, independently of p53 [73]. Furthermore, p73 and p63, but not p53, might activate the Notch signalling pathway and thus influence developmental cell fate decisions [74].

All together, genes of the p53 family might promote terminal differentiation of renal epithelial cells in the CD system and eventually initiate segment-specific transcription factors and cofactors ensuring segmental identity rather than specifying a cell lineage or cell fate [66].



**Fig. 2.9 Suggested role of p53 family in terminal nephron differentiation**, taken from [66]. **A)** Terminal nephron differentiation requires p53 stabilization and enhanced DNA binding activity. In the nephrogenic zone Pax-2 represses transcription of p53. In the differentiation zone high levels of p53 suppress transcription of PCNA and induces KLF4 transcription, which in turn activates p21 expression and sustains cell cycle arrest [66]. **B)** Activation of the AT<sub>1</sub>R by angiotensin II mediates phosphorylation of CREB, which allows interaction of CREB with p53 and the coactivator CBP, thus enhancing Bdkrb2 (B2R) transcription. CBP serves as a bridging molecule for the basal transcription machinery and as a histone acetyltransferase [66]. **C)** p53 and its homologue p73 are able to bind the promoters and activate the expression of renal function genes. However, the transcriptional activation requires additional transcription factors like KLF-4 or Foxi1 [66].

Pax-2 (paired box gene 2); PCNA (proliferating cell nuclear antigen); KLF4 (Krüppel-like factor 4); AT<sub>1</sub>R (angiotensin II receptor type 1); CREB (cAMP response-element binding protein); CBP (CREB-binding protein); Foxi1 (Forkhead box protein I1)

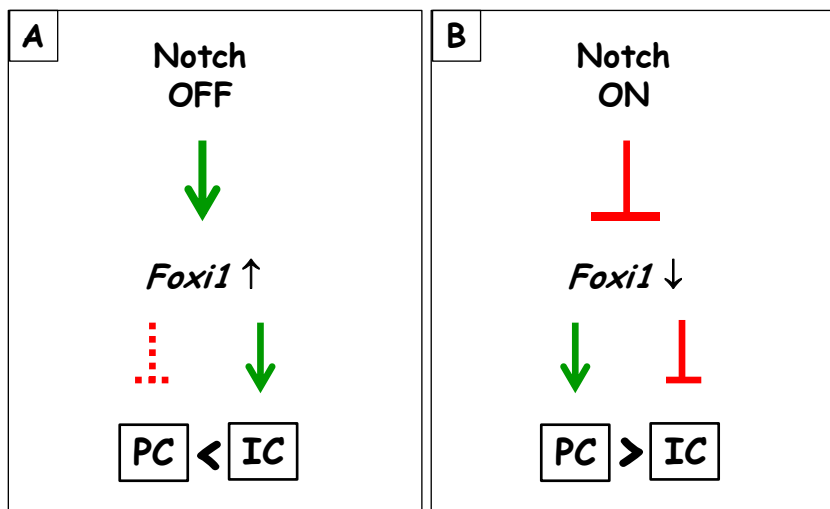
### **2.7.2 Notch signalling pathway**

The Notch signalling pathway is an evolutionary well conserved mechanism of intercellular crosstalk and a master regulator of development, cell fate determination, cell differentiation, tissue patterning, homeostasis and repair. The different actions of the Notch pathway can be broadly subdivided into three categories, which are 1) lateral inhibition, 2) lineage decisions, and 3) boundaries/inductive signalling [75]. Recent studies suggest Notch as a key regulator in renal development by establishing proximal epithelial fate during nephron segmentation and differentiation of principal cells in the renal CD system (Fig. 2.10) [76, 77].

This has been nicely demonstrated in a study by Jeong and colleagues [77]. In detail, the Notch signalling pathway was selectively disrupted in the renal CD system by a conditional KO of mind bomb 1 (Mib 1). Mib 1 is an E3 ubiquitin ligase that is necessary for endocytotic removal of Notch ligands and thus initiating Notch signalling [78]. However, inactivation of the Notch signalling pathway in the renal CD system, leads to nephrogenic diabetes insipidus due to reduced abundance of Aqp2 positive principal cells and their concomitant replacement by V-ATPase positive intercalated cells (Fig. 2.10 A) [77]. Interestingly, *Foxi1* mRNA abundance (see Chapter 2.7.3) was markedly increased in kidneys with inactivated Notch signalling. Intriguingly, constitutive activation of Notch signalling completely reverses the renal CD phenotype by restoring Aqp2 positive principal cells, virtually replacing all types of V-ATPase positive intercalated cells (Fig. 2.10 B). The constitutive activation of Notch was achieved by the introduction of the Notch intracellular domain (NCID) into the conditional Mib 1 deficiency phenotype [77]. Thus, Notch signalling critically influences the cellular heterogeneity of the mouse renal CD system.



Consistently, the Notch signalling pathway controls the mosaic pattern of multi-cilia cells and principal cells in the zebrafish pronephros by lateral inhibition [76, 79]. Furthermore, the Notch regulatory mechanism and members of the grainyhead family (see Chapter 2.7.4) most likely contribute to cell lineage determination of PSCs (proton-secreting cells) in *Xenopus laevis* and zebrafish larval skin, acting upstream of *foxi1* (Fig. 2.7, 2.8) [51].

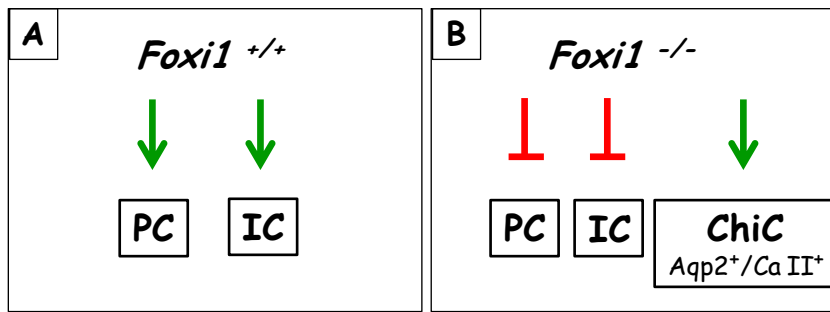


**Fig. 2.10 The Notch signalling pathway determines terminal differentiation of cells in the mammalian renal collecting duct system** (after [77]). **A)** Disruption of the Notch signalling pathway goes along with increased *Foxi1* mRNA expression and favours the abundance of V-ATPase positive intercalated cells (ICs) on the expense of Aqp2 positive principal cells (PCs). **B)** Constitutive activation of the Notch signalling pathway is accompanied by downregulation of *Foxi1* mRNA and results in the presence of Aqp2 positive PCs with a complete loss of V-ATPase positive ICs.

### **2.7.3 *Foxi1* deficiency**

Transcription factors are essential regulatory proteins in embryogenesis and organ development. There is a large family of forkhead box (FOX) transcription factors identified, which share homology in the forkhead box domain, a well conserved DNA binding motif of 80-100 amino acids, also known as the winged helix domain. At present, 50 FOX genes in the human genome and 44 Fox gene orthologues in mouse are identified and distinguished in 19 clades [80]. Based on phylogenetic criteria, a common nomenclature using alphabetic letters for distinct Fox classes (FOXA – FOXS) is agreed. The FOX transcription factors play important roles in embryo and organ development, cell cycle control, growth, differentiation, longevity and cancerogenesis in a variety of distinct species [81].

At least two Fox genes are involved in early kidney development, *Foxd1* and *Foxc1* [10]. Another member *Foxi1* was found to be expressed in distal tubules of fetal kidneys of later developmental stages and in the adult kidney [82], particularly in ICs of the renal CDs [83]. Furthermore, *Foxi1* expression was detected in the mouse inner ear and epididymis [84-86]. Analysis of the *Foxi1* consensus binding sequence identified potential target genes expressed in the renal distal tubule, among them are *Pds*, *Ae1*, *Ae4* and the *a4* and *B1 subunits of the V-ATPase* [82, 83, 85, 87]. Consistently, mice deficient for *Foxi1* develop distal renal tubular acidosis (dRTA), most evidently due to the loss of acid-base regulating proteins in renal ICs, such as *Pds*, *Ae1*, *Ae4* and the *B1 subunit of the V-ATPase* [83]. Interestingly, renal collecting ducts of *Foxi1* KO mice are composed of only a single cell type that is equally positive for the PC marker *Aqp2* and the IC marker *CaII*, instead of distinct populations of ICs and PCs (Fig. 2.11). Likewise, ultrastructural analysis of the CD system confirmed a lack of normal ICs in *Foxi1* null mutants. Instead of, a cell type neither displaying clear features of ICs nor strictly PC characteristics was found. Hence, all these data suggest *Foxi1* as a critical factor regulating the correct and functional patterning of the renal CD epithelium. Whether *Foxi1* is a direct or indirect regulator is still not known.

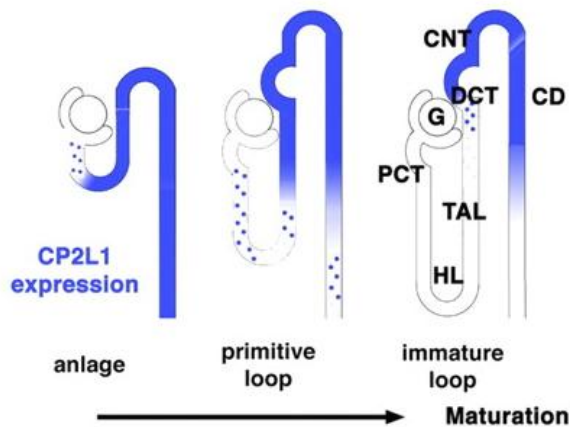


**Fig. 2.11 Relationship of *Foxi1* and terminal differentiation of mouse renal collecting duct cells**

(after [83]). **A)** The renal collecting duct (CD) of healthy mice consists of principal cells (PCs) and types of intercalated cells (ICs). **B)** However, PCs and ICs are absent in the renal CD system of *Foxi1* deficient mice and replaced by a chimeric cell (ChiC) type, coexpressing aquaporin 2 (Aqp2) and carbonic anhydrase II (CaII).

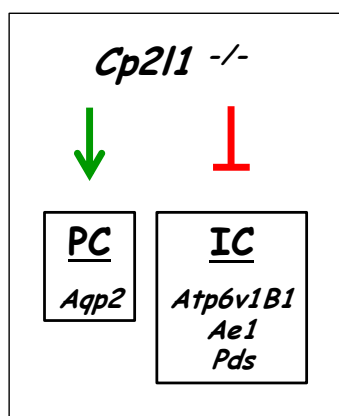
#### 2.7.4 *Cp2l1* deficiency

The family of LSF (late SV40 factor), also referred to as CP2 (CCAAT-binding protein 2 on  $\alpha$ -globin) related proteins comprises an ancient and evolutionary well conserved family of transcription factors which are rather ubiquitously expressed and usually transcriptional activators [88]. The LSF family consists of two branches, the LSF/CP2 subfamily and the grainyhead (GRH) subfamily. In the human and mouse genome three LSF-subfamily genes are identified, of which one is LBP-9/CRTR-1 (Leader binding protein 9, CP2-related transcriptional repressor-1) also known as TCFP2L1/CP2L1 (Transcription factor CP2-like1). CP2L1 is exceptional among the LSF-subfamily, as it is rather expressed in specific tissue and acts as a general repressor instead of activator [89]. In general, members of the grainyhead gene family play fundamental roles in the development and morphogenesis of epithelial tissues in a wide variety of species. Murine *Cp2l1* is characteristically expressed in the developing duct epithelium of exocrine glands including the salivary glands and in kidney [89, 90]. In detail, spatio-temporal *Cp2l1* expression in the fetal mouse kidney is differentially regulated in both, the nephrogenic and ureteric bud derived epithelia (Fig. 2.12). However, in the postnatal maturing kidney, *Cp2l1* expression becomes restricted to the DCT, CNT, CCD and OMCD.



**Fig. 2.12 Differential expression of Cp2l1 in the developing mouse kidney,** taken from [90].

Developmental maturation of functional salivary gland and renal distal tubule epithelium was defective in Cp2l1 deficient mouse embryos [90]. In particular, Cp2l1 depleted kidneys lack the expression of *Clcnkb*, *Gk-6* and *Fxyd2c*, genes which are characteristically present in the TAL, CNT and DCT, respectively. Moreover, mRNA expression of *Atp6v1b1*, *Ae1* and *Pds* was completely absent in Cp2l1 deficient kidneys (Fig. 2.13). Unfortunately, more detailed characterization of the renal CD morphology is missing, as well as data assessing the acid-base homeostasis in these mice, despite the observation of a mild urinary concentrating defect. Anyhow, Cp2l1 deficiency in mice correlates with incomplete renal CD differentiation, strongly affecting the characteristic structure and function of ICs. However, regulatory pathways and mechanisms involving transcriptional control of Cp2l1 are still missing.



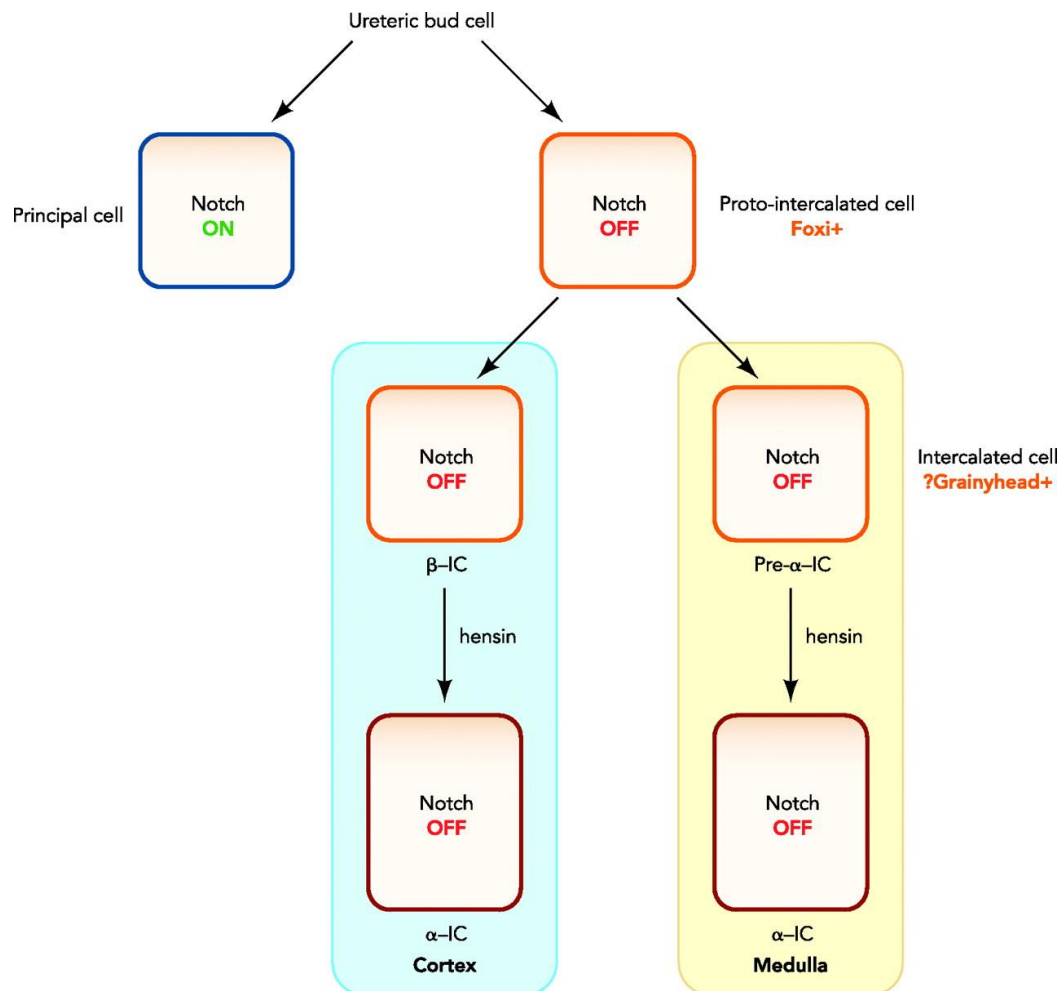
**Fig. 2.13 Correlation of Cp2l1 deficiency with the expression of genes usually present in mouse renal collecting duct cells** (after [90]). Loss of *Cp2l1* in kidneys of mice does not alter the mRNA expression of aquaporin 2 (*Aqp2*), a marker for principal cells (PCs). However, *Cp2l1* deficiency goes along with the absence of *Atp6v1B1*, *Ae1* and *Pds* mRNA, usually expressed by types of intercalated cells (ICs).

### **2.7.5 *Dmbt1 (Hensin) deficiency***

Besides the importance of regulatory transcription factors and signalling molecules, another group of proteins, namely integrins and extracellular matrix (ECM) proteins, play a role in terminal differentiation of renal epithelial cells. Initially discovered in cultured epithelial cells derived from rabbit kidney, the ECM protein hensin is proposed crucial for terminal differentiation of renal intercalated cells and the hypothesized transdifferentiation of types of ICs upon acid-base disturbances (chapter 5.2) [91-93]. Hensin is expressed during embryonic development and with species differences in the expression levels in pancreas, intestine, gallbladder, salivary glands and nasal/olfactory/vomerolnasal epithelium [94]. It has been described previously by different investigators and thus is known, beside other names, also as *Dmbt1* (deleted in malignant brain tumors 1) or muclin (mucin-like glycoprotein). However, it has been reported, that hensin is synthesized and secreted to the extracellular matrix as a monomer or polymer. The polymerization of hensin seems to mediate interconversion of type B-ICs to type A-ICs and requires the presence of at least 3 additional proteins (galectin 3, cis-trans peptidyl prolyl isomerase and integrin  $\alpha v \beta 1$ ) [95-98].

Recently, a mouse model with conditional deletion of hensin from renal intercalated cells was investigated [99]. The mice were reported to lack renal type A-ICs all along the CD system and consequently develop complete dRTA due to renal bicarbonate wasting. Essentially, type B-ICs were preserved in the renal cortex of the mutant mice, while a novel IC phenotype was discovered in the inner medulla. Here, the authors found an uncertain type of ICs, which displays the ultrastructural characteristics of type B-ICs but lacks pendrin expression. This novel cell type is suggested to be the progenitor of medullary type A-ICs. In the same study, another mouse model with a conditional deletion of integrin  $\beta 1$  from renal ICs was compared, since integrin  $\beta 1$  was suggested critical for hensin polymerization and deposition to the ECM. Accordingly, a phenotype similar to that of hensin deficient mice was observed. However, despite intensive studies, a putative molecular mechanism of hensin mediating renal IC

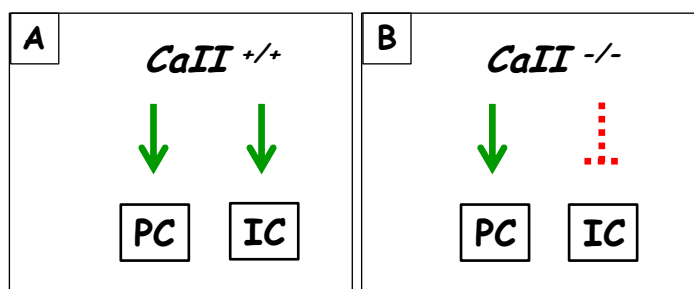
differentiation has never been suggested. Nevertheless, hensin was very recently integrated into an oversimplified schematic model of intercalated cell differentiation (Fig. 2.14) [100].



**Fig. 2.14 Proposed model for renal intercalated cell differentiation**, taken from [100].

### 2.7.6 Carbonic anhydrase II deficiency

Besides transcription factors and signalling molecules the presence of the enzyme carbonic anhydrase II (CaII) was found to be a critical determinant of terminal IC differentiation. Renal ICs express high levels of the cytosolic enzyme CaII which catalyzes the rapid conversion of CO<sub>2</sub> and water to H<sub>2</sub>CO<sub>3</sub>. CaII deficiency in mouse and men is known to cause distal renal tubular acidosis (dRTA). Moreover, ICs of the CCD and particularly MCD are severely reduced and replaced by PCs in kidneys of CaII deficient mice (Fig. 2.15) [101, 102]. Another study showed that the phenomenon of IC depletion in CaII deficient mice develops during late postnatal maturation of the kidney [103]. In detail, a remodelling of the MCD favouring PC abundance on the expense of ICs was only detectable after 3 weeks of life. These observations suggest that the presence or absence of CaII directly influences the cell type diversity in the renal CD.



**Fig. 2.15 Carbonic anhydrase II (CaII) deficiency leads to depletion of intercalated cells (ICs) in mouse kidney** (after [101-103]). Detailed description in text.

Notably, chronic pharmacologic inhibition of CaII with acetazolamide in rats was shown to induce distinct remodelling of the renal CD system [104]. Here, type A-IC abundance in the renal CCD was increased with a corresponding decrease in type B-ICs, without affecting the ratio of total IC to PC. Furthermore, the relative number of type A-ICs in the outer medulla was increased on the expense of PCs. In addition, the size of ICs in CCD and OMCD was markedly increased. In contrast, A-ICs were strongly reduced in the inner medulla of acetazolamide treated rats, but not completely absent as it was reported for CaII deficient mice. These data are consistent with the concept of adaptive nephron plasticity (see Chapter 5.2.1).

### 3. RENAL ACID-BASE TRANSPORT

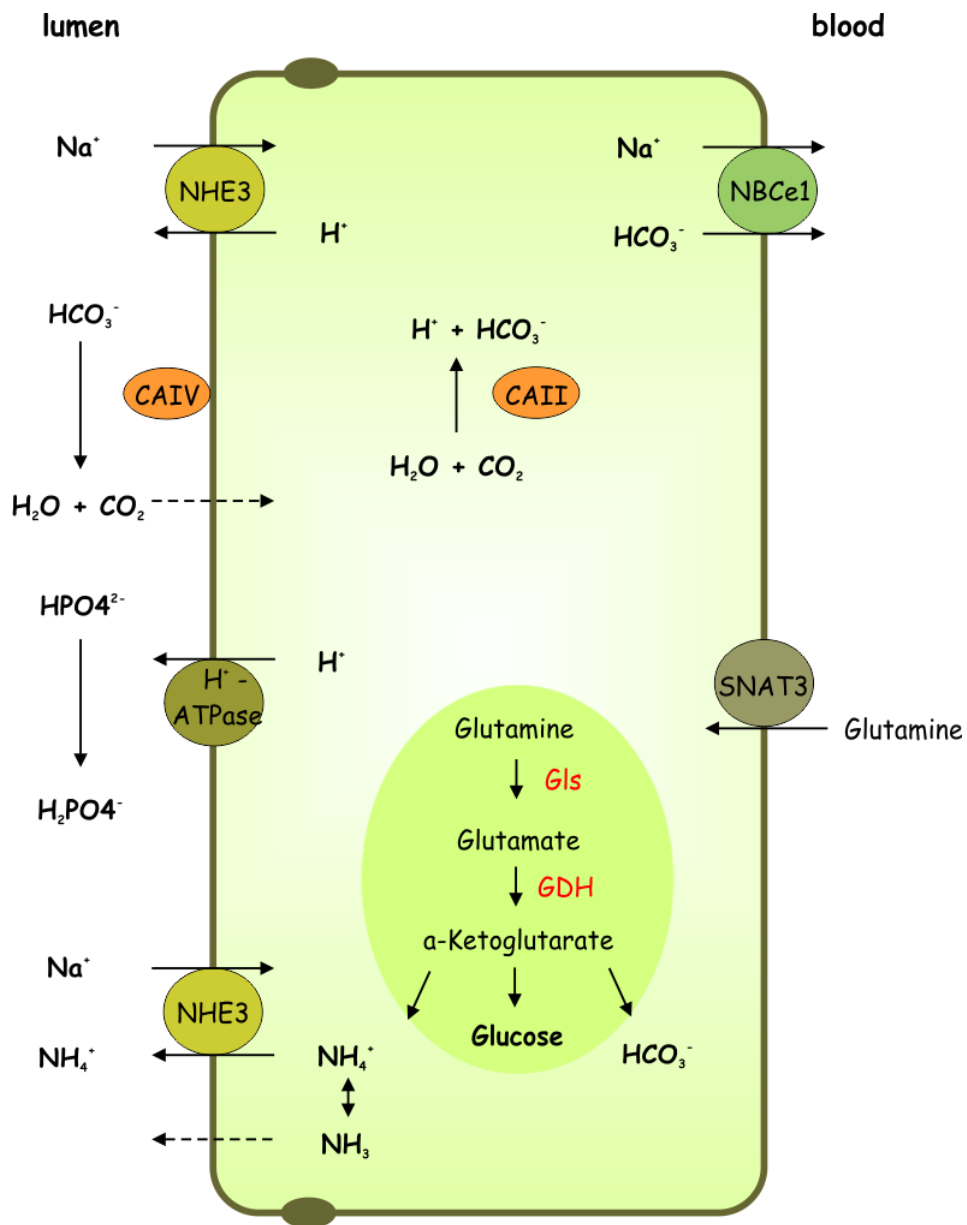
The kidneys ability to produce acidic urine is an important mechanism to adapt to intermediate and chronic imbalances in systemic acid-base homeostasis. Renal excretion of acids is enabled by three regulatory pathways along the nephron: 1) reabsorption of filtered  $\text{HCO}_3^-$  in the proximal tubule, 2) ammoniagenesis and excretion of ammonia and titratable acids, and 3) excretion of excess protons in the distal nephron.

#### 3.1 *Renal bicarbonate handling*

Bicarbonate in the blood is freely filtered in the renal glomerulum and thus needs to be recovered from the renal tubular lumen to prevent loss into the urine. The proximal tubule is the main site of renal  $\text{HCO}_3^-$  reabsorption, accounting for 80% [105, 106]. As illustrated in Fig. 3.1, luminal  $\text{HCO}_3^-$  and  $\text{H}^+$  are converted to  $\text{CO}_2$  and  $\text{H}_2\text{O}$  by the action of the membrane bound carbonic anhydrase IV (CAIV). Carbon dioxide enters the proximal tubule cell rapidly by diffusion where it is once more combined with  $\text{H}_2\text{O}$  to form  $\text{HCO}_3^-$  and  $\text{H}^+$ . This intracellular hydration is catalyzed by carbonic anhydrase II (CAII), which is a soluble isoform. While the  $\text{H}^+$  exits the cell to the luminal site, mostly in exchange for  $\text{Na}^+$ , via the  $\text{Na}^+/\text{H}^+$  exchanger NHE3, the formed  $\text{HCO}_3^-$  is cotransported together with  $\text{Na}^+$  to the blood by the basolateral  $\text{Na}^+/\text{HCO}_3^-$  cotransporter NBCe1.

The remaining 20 % of luminal  $\text{HCO}_3^-$  are finally reabsorbed in the thick ascending limb of the loop of Henle [107] and the distal tubule [108].



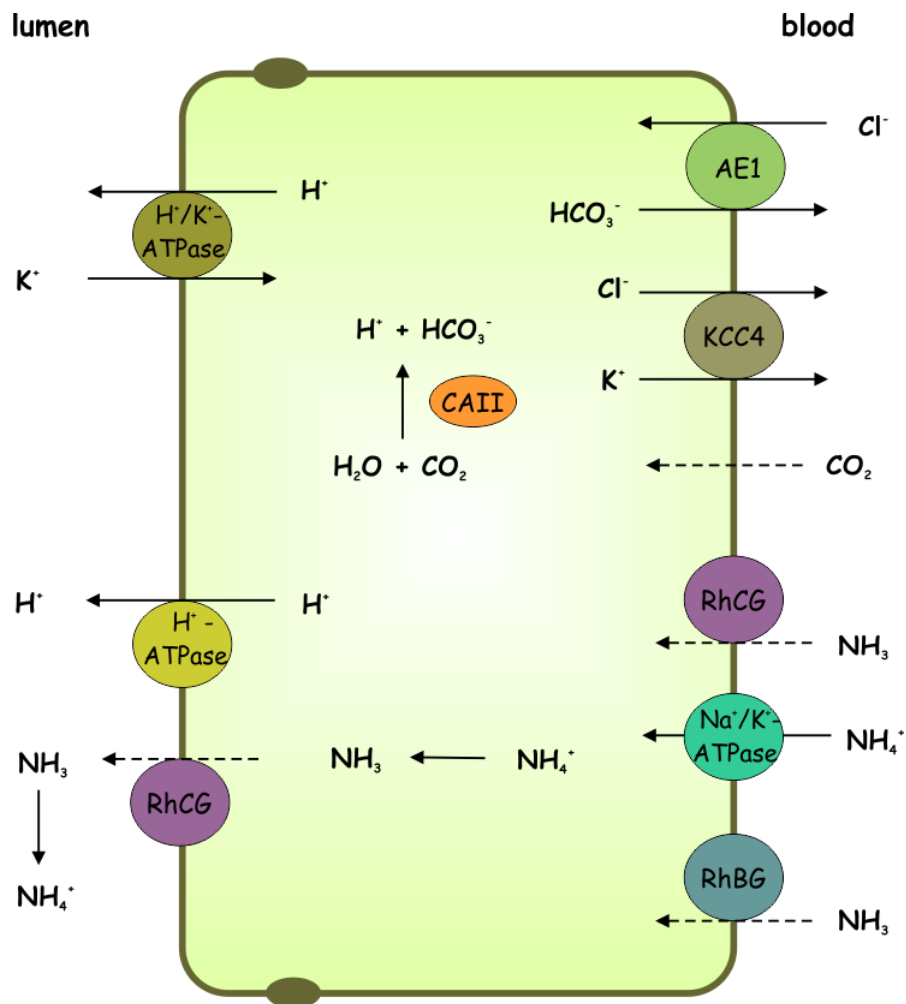


**Fig. 3.1 Bicarbonate reabsorption in proximal tubule** (adapted after [1]), detailed description in text.

### **3.2 Ammoniogenesis**

Beside the recovery of bicarbonate from the urine, the kidneys are able to generate new bicarbonate from the synthesis of ammonium ( $\text{NH}_4^+$ ). Ammonium is primarily produced from glutamine in mitochondria of the proximal tubule (Fig. 3.1) [109]. Glutamine, provided by the liver due to amino acid break down, enters the proximal tubule cells mostly by the  $\text{Na}^+$  coupled cotransporter SNAT3 [110]. Inside the mitochondria, glutamine is cleaved to glutamate and ammonium ( $\text{NH}_4^+$ ) by glutaminase. Glutamate is then further converted to  $\alpha$ -ketoglutarate and a second  $\text{NH}_4^+$  by glutamate dehydrogenase (GDH). Ammonium inside the cell dissociates into  $\text{H}^+$  and gaseous  $\text{NH}_3$ , which diffuses into the tubular lumen, where it binds again to  $\text{H}^+$  and is trapped as  $\text{NH}_4^+$ . The protons are consumed to form glucose from  $\alpha$ -ketoglutarate, thus generating bicarbonate. Ammonium might also be secreted into the lumen by the apical  $\text{Na}^+/\text{H}^+$  exchanger NHE3, replacing the  $\text{H}^+$  as the preferred substrate. Along the different parts of the loop of Henle (LOH), ammonium is secreted and primarily reabsorbed in the thick ascending limb (TAL). This leads to the accumulation of  $\text{NH}_4^+$  in the interstitium of the outer medulla, partially favouring non-ionic diffusion of  $\text{NH}_3$  into the lumen of the medullary collecting ducts, where it is excreted to the final urine along with  $\text{H}^+$ .

As recently discovered, ammonia secretion in the inner medullary collecting duct, is not only passive, but mediated by the  $\text{Na}^+/\text{K}^+$ -ATPase and the Rhesus glycoproteins RhBG and RhCG [111-114]. These specialized transporters are predominantly present in principal cells and type-A intercalated cells of the collecting ducts [112] (Fig. 3.2). Mice deficient for Rhcg display an incomplete form of distal renal tubular acidosis (dRTA), due to insufficient  $\text{NH}_3$  excretion to the urine, indicating its relevance (see chapter 7.1.2) [114].



**Fig. 3.2 Ammonia excretion in type-A intercalated cell of renal collecting duct** (adapted after [109]), detailed description in text.

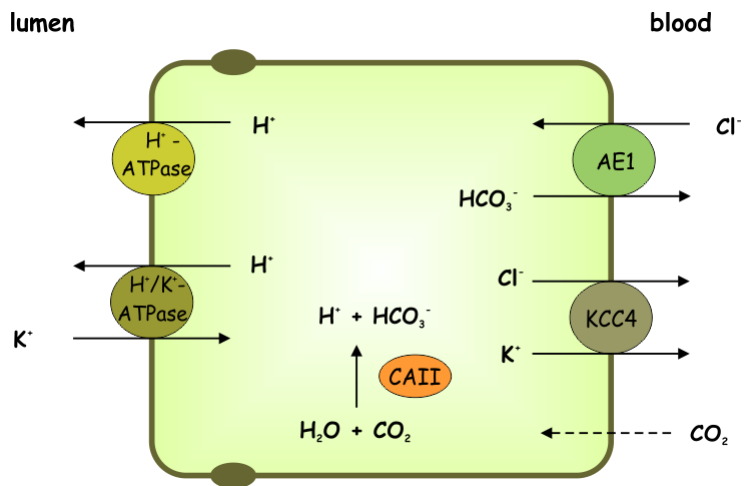
### **3.3 Renal use of titratable acids**

Protons excreted with the urine are not only bound as  $\text{NH}_4^+$ , but are also bound to other buffers, the so called titratable acids (TA). By definition, titratable acidity describes the amount of acid required to titrate the urine buffers from the plasma pH to urine pH [115]. The most important urinary titratable acids are phosphate (Fig. 3.1) and creatinine. However, other organic salts such as sulfate, citrate, lactate, pyruvate and urate also contribute to urinary buffering and thus, titratable acidity. The term net acid excretion (NAE) describes the sum of titratable acids and  $\text{NH}_4^+$  minus the urinary  $\text{HCO}_3^-$  concentration in the urine (Eqn. 3) [1].

Eqn. 3       $\text{NAE} = (\text{TA} + \text{NH}_4^+) - \text{HCO}_3^-$

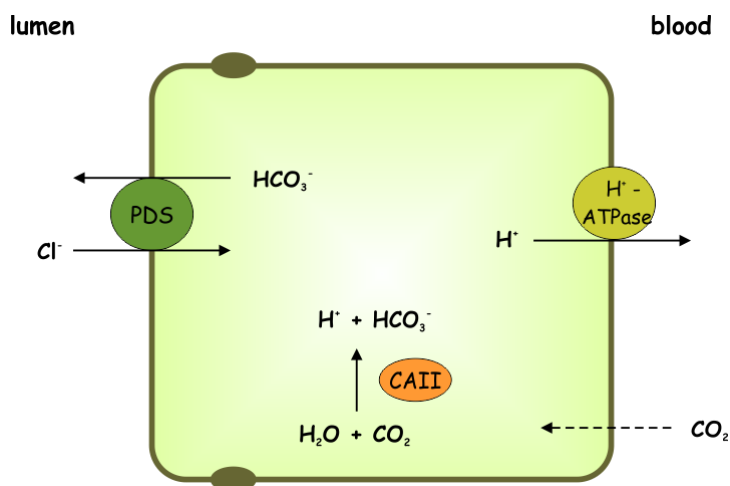
### **3.4 Acid-base transport in the distal nephron**

While the proximal tubule plays a major role in the *de novo* generation and reabsorption of bicarbonate, the distal tubule and the collecting ducts are the site of urinary fine tuning and final acidification [116]. Today, the distal convoluted tubule (DCT), the connecting tubule (CNT) and the initial collecting tubule (ICT) are taken together as the classical distal tubule, even though they display significant cell heterogeneity. The collecting ducts (CD) are distinguished in cortical collecting ducts (CCD), outer medullary collecting ducts (OMCD) and inner medullary collecting ducts (IMCD). As already described, the collecting ducts are comprised of at least three different cell types, principal cells, type-A and type-B intercalated cells. The principal cells are responsible for salt and water homeostasis. Type-A intercalated cells are acid secretory cells, since they secrete protons into the urine and transport bicarbonate into the blood. A-ICs are characterized by the luminal expression of the V-ATPase and basolateral expression of the  $\text{Cl}^-/\text{HCO}_3^-$  exchanger AE1 (SLC4A1, Band 3) (Fig. 2.6, 3.3).



**Fig. 3.3 Model of type-A intercalated cell (A-IC) without NH<sub>3</sub>/NH<sub>4</sub><sup>+</sup> pathway**, adapted from [116], detailed description in text.

Type-B intercalated cells (B-ICs) are bicarbonate secreting cells and show opposite features of type A-ICs with the luminal expression of the Cl<sup>-</sup>/HCO<sub>3</sub><sup>-</sup> exchanger pendrin (SLC26A4, PDS) and the basolateral presence of V-ATPase (Fig. 2.6, 3.4).



**Fig. 3.4 Model of type-B intercalated cell (B-IC)**, adapted from [116], detailed description in text.

However, proton and bicarbonate excretion by the collecting ducts is strongly dependent on the presence of NH<sub>3</sub> and titratable acids in the urine and is influenced by different factors, such as systemic acid-base and electrolyte status, several hormones and nutrition [116, 117].

#### 4. DISTURBANCES OF ACID-BASE HOMEOSTASIS

Systemic and renal acid-base regulation is influenced by a multitude of physiologic and pathophysiologic factors including respiration, nutrition and physical activity, as well as medication and various epithelial transporter disorders. Clinically, acidosis defines a blood pH lower than 7.35, while alkalosis exists with a blood pH higher than 7.45. Typically, four basal pH disturbances of different origin (Tab. 4.1) are distinguished: 1) respiratory acidosis, 2) respiratory alkalosis, 3) metabolic acidosis, and 4) metabolic alkalosis. In general, respiratory derived acid-base disturbances are primarily caused by alterations in blood  $p\text{CO}_2$  due to inappropriate pulmonary ventilation. In contrast, metabolically derived pH imbalances are usually caused by accumulation or loss of non-volatile acids and therefore changes in blood  $\text{HCO}_3^-$  concentration. Furthermore, metabolically derived pH disturbances may be distinguished in extrarenal and renal origin. Extrarenal metabolic pH disturbances might be caused by nutrition (i.e. high protein content), diarrhoea and excessive sports. However, systemic pH imbalances might also arise because of renal tubular dysfunction. In any case, the body tries to restore systemic pH as close to normal by acute and long-term regulatory mechanisms.

**Tab. 4.1 Clinical types of acid-base disturbances**, modified after [1]

Disturbance	Symptom	Response
Respiratory acidosis	$\uparrow p\text{CO}_2$	$\uparrow \text{HCO}_3^-$
Respiratory alkalosis	$\downarrow p\text{CO}_2$	$\downarrow \text{HCO}_3^-$
Metabolic acidosis	$\downarrow \text{HCO}_3^-$	$\downarrow p\text{CO}_2$
Metabolic alkalosis	$\uparrow \text{HCO}_3^-$	$\uparrow p\text{CO}_2$

Clinically, determination of the serum anion gap helps to identify the cause of metabolic acidosis [118]. The anion gap is a calculated value, given by subtraction of the major anions [ $\text{HCO}_3^-$ ] and [ $\text{Cl}^-$ ] from the major cation [ $\text{Na}^+$ ] (Eqn. 4).

$$\text{Eqn. 4} \quad \text{Anion Gap} = [\text{Na}^+] - ([\text{HCO}_3^-] + [\text{Cl}^-])$$

Thus, the anion gap is usually a positive value ( $12 \pm 4$  meq/l), reflecting the quantity of anions which are not balanced by cations [119]. The result of the anion gap is classified as high or normal and allows the following interpretations. A high anion gap acidosis may be observed in the presence of additional metabolic acids consuming bicarbonate (i.e. lactic acidosis, ketoacids) or in renal failure, when the kidneys do not sufficiently reabsorb bicarbonate or secrete acids. Normal anion gap acidosis, also called hyperchloremic acidosis, goes along with the loss of bicarbonate compensated by an equivalent increase in serum chloride concentration. This form is usually caused by renal tubular acidosis, diarrhoea and ingestion of ammoniumchloride or hydrochloric acid.

#### **4.1 Respiratory acidosis**

Respiratory acidosis occurs during insufficient alveolar ventilation (hypoventilation) and leads to increased arterial  $\text{pCO}_2$  levels (hypercapnia) which is initially corrected by  $\text{HCO}_3^-$  buffering from the extracellular fluid. On the long term, the kidneys need to correct the distorted blood [ $\text{HCO}_3^-/\text{CO}_2$ ] by reabsorbing all filtered  $\text{HCO}_3^-$  and enhancing the generation and secretion of all available acids by upregulating renal acid-base transporters [120, 121].

## **4.2    *Respiratory alkalosis***

Hypocapnia describes a state of low arterial  $p\text{CO}_2$  levels caused by alveolar hyperventilation. This leads to respiratory alkalosis and fall in plasma  $[\text{HCO}_3^-]$ . Chronic hypocapnia further decreases extracellular fluid  $[\text{HCO}_3^-]$  as a result of reduced renal net acid excretion [120, 121].

## **4.3    *Metabolic acidosis***

Metabolic acidosis is a severe acid-base imbalance that may be caused by numerous factors, such as starvation, diarrhoea, intoxication or disease complications like diabetic ketoacidosis. Acute metabolic acidosis, lasting up to several days, is much more common than chronic forms [122]. However, the immediate compensatory response to metabolic acidosis is an increase in alveolar ventilation, thus exhaling more  $\text{CO}_2$ . This is usually achieved by enhancing the respiratory tidal volume and frequency. However, the final correction of plasma  $[\text{HCO}_3^-/\text{CO}_2]$  needs to be established by the kidney. Thus, renal excretion of  $\text{H}^+$  and  $\text{HCO}_3^-$  reabsorption is maximally upregulated. Additionally, ammoniagenesis and final excretion of  $\text{NH}_4^+$  increases markedly and becomes the main  $\text{H}^+$  extrusion pathway. Detailed adaptive responses of the kidney to acid loads will be discussed later on (chapter 5).



#### **4.4    *Metabolic alkalosis***

Two conditions may lead to metabolic alkalosis, namely the loss of hydrogen ions (i.e. by vomiting) or the inappropriate increase in plasma  $[\text{HCO}_3^-]$  (i.e. abuse of steroids and diuretics). Likewise to metabolic acidosis, the respiratory system responds and adapts immediately to metabolically derived alkalosis. Here, the respiratory frequency slows down significantly, leading to the accumulation of  $\text{pCO}_2$  in the blood. Again, the kidney is responsible for the final and persistent correction. During metabolic alkalosis, acid secretion and bicarbonate reabsorption by the kidney is markedly reduced. Moreover, type B-ICs of the renal cortical collecting ducts may even switch from net  $\text{H}^+$  secretion to  $\text{HCO}_3^-$  secretion [1, 123].

#### **4.5    *Renal tubular acidosis***

Impaired renal acidification leads to renal tubular acidosis (RTA), a form of metabolic acidosis with hyperchloremic anion gap in the setting of normal glomerular filtration [124]. Generally, acquired or inherited RTA may develop either because of insufficient proximal bicarbonate reabsorption or due to inadequate distal urinary acidification. However, based on the clinical symptoms and the underlying functional defect, four different types of RTA are distinguished: type 1 classic distal (dRTA), type 2 proximal (pRTA), type 3 combined proximal and distal RTA, and type 4 distal-like hypoaldosteronism (hyperkalemic dRTA) [125, 126].

#### **4.5.1 Type 1 RTA**

Distal renal tubular acidosis (dRTA) is characterised by the renal inability to acidify the urine, leading to metabolic acidosis with variable severity of accompanying hypokalemia, hypercalciuria and hypocitraturia. The underlying defect is usually an acquired or inherited form of impaired acid secretion by type A-ICs of the renal collecting ducts. The impact of inadequate urine acidification might result in spontaneous systemic metabolic acidosis (complete dRTA) or metabolic acidosis after an imposed acid load (incomplete dRTA) [127]. Therapeutically, adequate bicarbonate supplementation is sufficient to restore plasma bicarbonate to near normal levels. The oral application of potassium citrate is favoured over sodium bicarbonate, as it simultaneously treats the accompanying and often life-threatening hypokalemia [128].

Primary familial dRTA is caused by mutations of epithelial transporters essentially expressed in renal type A-ICs. In dependence of the mutation and the affected transporter respectively, autosomal- dominant and –recessive forms of dRTA are known (Tab. 4.2) [126, 127]. Most commonly, loss of function mutations of the  $\text{Cl}^-/\text{HCO}_3^-$  exchanger AE1 or the V-ATPase were identified in patients with primary familial dRTA [129-134].

However, dRTA frequently also develops as a secondary disorder, often in consequence of autoimmune diseases such as Sjögren`s syndrome [135-138].

**Tab. 4.2 Subtypes and features of inherited dRTA**, adapted from [126].

<b>dRTA Inheritance</b>	<b>Age at Presentation</b>	<b>Clinical Features</b>	<b>Transporter</b>
Dominant	Adulthood	mild/compensated Metabolic Acidosis Hypokalemia (variable) Hypercalciuria Hypocitraturia Nephrolithiasis Nephrocalcinosis sometimes Rickets/ Osteomalacia Secondary Erythrocytosis	AE1 (SLC4A1)
Recessive	Childhood	Metabolic acidosis with Hemolytic Anemia	AE1 (SLC4A1)
Recessive	Infancy/ Childhood	early onset bilateral sensorineural hearing loss Metabolic Acidosis early Nephrocalcinosis growth retardation Rickets	B1 subunit of V-ATPase (ATP6V1B1)
Recessive	Infancy/ Childhood	as above with later onset of hearing loss or no deafness	a4 subunit of V-ATPase (ATP6V0A4)

#### **4.5.2 Type 2 RTA**

Proximal renal tubular acidosis (pRTA) is usually caused by defective proximal tubule bicarbonate reabsorption with intact distal acidification. This leads to excessive loss of bicarbonate by the kidney and subsequent metabolic acidosis with hypokalemia [139]. Primary isolated pRTA is a rare disorder and associated with dominant and recessive inherited mutations in the  $\text{Na}^+/\text{HCO}_3^-$  cotransporter NBCe1 (SLC4A4) (Fig. 3.1). Since NBCe1 is also expressed in the eye, pRTA might come along with ocular abnormalities [140].

#### **4.5.3 Type 3 RTA**

Type 3 RTA is caused by carbonic anhydrase II (CAII) deficiency and combines proximal and distal RTA features. It is characterized by reduced proximal bicarbonate reabsorption and distal inability to acidify the urine despite systemic acidosis [141]. CAII plays an important role in bone and central nervous system (CNS) metabolism. Therefore, CAII malfunction frequently leads to osteopetrosis, cerebral calcification, and mental retardation [142, 143].

#### **4.5.4 Type 4 RTA**

Type 4 RTA is actually a primary disorder in renal electrolyte handling with secondary normal anion gap acidosis. In general, renal aldosterone deficiency or resistance are thought to be the underlying defect. Thus, hyperkalemia and hyperchloremia are unique characteristics of type 4 RTA. The mild metabolic acidosis develops secondary to the hyperkalemia, which induces an intracellular alkalosis impairing renal proximal  $\text{NH}_4^+$  generation and excretion [144].

## **5. RENAL ADAPTATION TO SYSTEMIC ACIDOSIS**

Systemic acidosis requires compensatory responses of the kidney function increasing the capacity of acid-base and electrolyte handling with cellular reorganization and even remodelling of the nephron [145-148]. Furthermore, the body has other mechanisms that contribute to renal compensatory regulation of acid-base imbalance, such as increase of glomerular filtration rate (GFR) [1], hormones that directly stimulate kidney function (angiotensin II, endothelin) [149, 150] and hormones (i.e. parathyroid hormone, vitamin D) affecting bone resorption and thus recruitment of additional non-carbonate buffers [151, 152].

### ***5.1 Adaptation of proximal renal tubule***

In response to metabolic acidosis the renal proximal tubule undergoes a set of adaptations to maximally recover filtered bicarbonate [153] and generate new bicarbonate to buffer excess systemic acids. Increased acid loads enhance the capacity of proximal bicarbonate reabsorption by modulating the activity and/or abundance of apical and basolateral transport mechanisms [154]. The apically located  $\text{Na}^+/\text{H}^+$  exchanger NHE3 plays a crucial role in transcellular bicarbonate reabsorption (Fig. 3.1) and enhanced transport activity, as well as increase in its abundance in the proximal tubule and TAL during metabolic acidosis have been demonstrated [154-157]. Furthermore, the V-ATPase is an important apical  $\text{H}^+$  extrusion pathway, accounting for up to 40% of sodium independent bicarbonate transport, which is rapidly regulated by  $\text{pCO}_2$  and pH [158, 159]. Moreover, acidic pH modulates basolateral  $\text{Na}^+/\text{HCO}_3^-$  cotransport mediated by NBCe1 and increases the abundance of transporter at the cell membrane [160-163]. In turn, mutations of NBCe1 lead to proximal renal tubular acidosis (chapter 4.5.2) [164, 165].

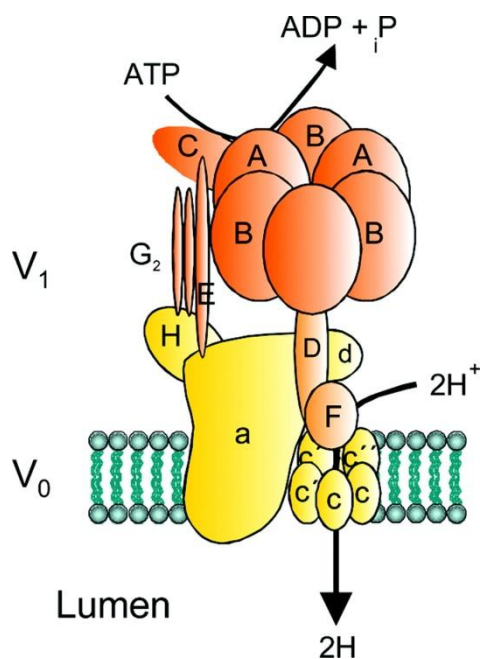
However, effective renal acid buffering requires *de novo* generation of additional bicarbonate by the synthesis and excretion of ammonium and excess acids. Ammonium is enzymatically generated from the amino acid glutamine in renal proximal tubule cells (Fig. 3.1). During metabolic acidosis, the excretion and synthesis of ammonium is highly upregulated [166] involving the supply and uptake of glutamine into renal tubular cells [167].

Additionally, the kidney increases the excretion of titratable acids, mainly phosphate (Fig. 3.1). In the proximal tubule, two unrelated gene families (SLC20 and SLC34) of phosphate transporters are expressed [168, 169]. Transport activity and abundance of members of the SLC34 family were found to be regulated upon acid-base disturbances [170]. In response to metabolic acidosis elevated levels of phosphate are excreted into the urine, where it combines with excess protons acting as a titratable acid. Since elevated circulating phosphate is partially released from bone mass, chronic systemic acidosis may lead to osteomalacia or osteopetrosis, as well as nephrocalcinosis and nephrolithiasis [171].

## **5.2 Adaptation of distal renal tubule**

The renal response to metabolic acidosis does not only require adaptations of the proximal tubule, but also of the distal nephron. The importance of the final urinary acidification and fine tuning by the distal nephron is also highlighted by the dysfunction of numerous transporters leading to dRTA. Within the collecting duct, the vacuolar type  $H^+$ -ATPase (V-ATPase) is the major protein responsible for  $H^+$  secretion into urine. It is differentially expressed in all types of intercalated cells and thus involved in mediating renal net  $H^+$  secretion and  $HCO_3^-$  generation or  $H^+$  reabsorption and net  $HCO_3^-$  secretion (Fig. 2.6) [133]. The V-ATPase is a heteromultimeric complex, consisting of 13 different subunits which are assembled in two domains (Fig. 5.1) [172]. Some of the identified subunits are almost exclusively expressed in the kidney among them are the a4 (ATP6V0A4) and B1 (ATP6V1B1)

subunits. As indicated in Fig. 2.6, 3.3 and 3.4, the plasma membrane location of the V-ATPase in intercalated cells (IC) is strongly dependent on the type of IC. Interestingly, acid-base disturbances do not seem to alter the mRNA or protein expression levels of renal V-ATPases, but trigger its redistribution within the cell [40]. While acidosis promotes the insertion of the proton pump into the cell membrane from a subplasma membrane pool of V-ATPase vesicles, alkalosis drives the retrieval of V-ATPase from the membrane into the submembrane vesicular pool [147]. In addition, acid-base disturbances regulate the protein abundance of the  $\text{Cl}^-/\text{HCO}_3^-$  exchangers Ae1 and Pds (Fig. 2.6). Acid-loads were shown to increase the protein abundance of Ae1 and decreases Pds, whereas alkali-loads decrease Ae1 abundance and increase Pds [173].



**Fig. 5.1 Structural model of vacuolar type  $\text{H}^+$ -ATPase (V-ATPase),** taken from [172].

### **5.2.1 Adaptive plasticity of distal renal tubule**

In addition to acid-base related changes of the cellular protein composition and localization, acute and especially long-term acid-base imbalances are accompanied by changes in the relative number and axial distribution of renal type A-ICs and B-ICs [104, 145, 146]. It has been observed that the relative number of type A-ICs increases dramatically in rodents receiving chronic oral acid loads. Strikingly, the relative number of type B-ICs decreases at the same time, while the ratio of total ICs to PCs is not changed [173, 174]. The opposite phenomenon is true for rodents receiving chronic alkali loads, since the relative number of type B-ICs increases on the expense of type A-ICs. These empiric data raised several hypotheses, which are:

- 1) IC progenitors might exist in the CD epithelium and become activated upon acid-base perturbations [175-177].
- 2) Terminally differentiated ICs proliferate or become apoptotic upon prolonged functional request [138, 146, 178, 179].
- 3) The concept of the plasticity of functional epithelial polarity was raised, a process where type B-ICs might convert into type A-ICs or *vice versa*. Moreover, type B-ICs are postulated precursors of type A-ICs [91, 174, 180].

Since then, several studies have been performed testing either idea. Accordingly, circulating (i.e. haematopoietic stem and progenitor cells, HSPCs) or tissue specific residual progenitor cells giving rise to new differentiated renal epithelial cells seem rather important for postinjury repair. Furthermore, the notion of proliferation or apoptosis of terminally differentiated cells upon exogenous stimuli (i.e. acid-base or electrolyte disturbances) is more tempting than the concept of adaptational reversal of functional epithelial polarity. In fact, the latter theory was never proofed convincingly.



### **5.2.2 *Gdf15 is involved in early adaptive proliferation of renal type A-IC***

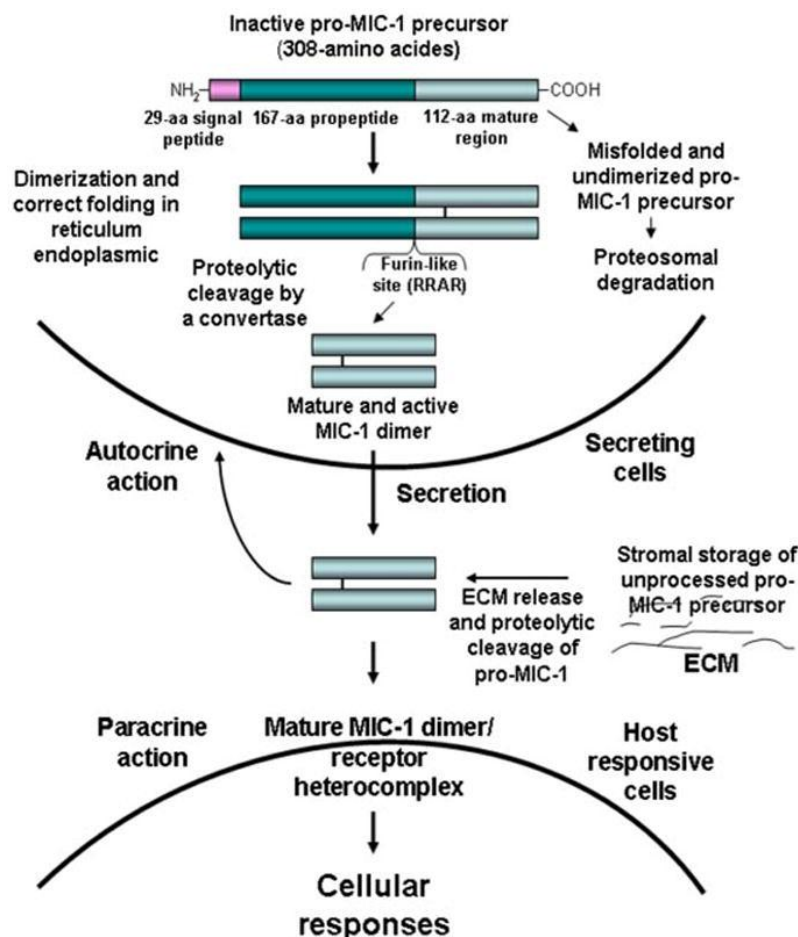
Recently, the growth differentiation factor 15 (Gdf15), a member of the transforming growth factor- $\beta$  (TGF- $\beta$ ) superfamily was found to be involved in the proliferation of type A-ICs during early adaptation of the kidney to metabolic acidosis [146]. The murine orthologue Gdf15 is also designated as macrophage inhibitory cytokine-1 (MIC-1), prostate-derived factor (PDF), placental TGF- $\beta$  (PTGF- $\beta$ ), placental bone morphogenetic protein (PLAB) and non-steroidal anti inflammatory drug-activated gene-1 (NAG-1), reflecting its pleiotropic functions [181]. As typically reported for TGF- $\beta$  superfamily members, Gdf15 is a multifunctional acting factor and involved in the control of embryonic development and pregnancy, osteogenic and haematopoietic development, cellular responses to stress stimuli, inflammation and tissue repair after acute injury.

During protein synthesis, Gdf15 undergoes extensive peptide processing leading to several manifestations, such as signal-peptide, propeptide, and mature peptide, which are associated with distinct molecular mechanisms (Fig. 5.2) [181]. The biological active mature protein is a secreted dimer, serving multiple autocrine and paracrine actions upon a multitude of stimuli (detailed review by [181]).

In the mature mouse kidney, low mRNA expression levels all along the nephron were reported under basal conditions [146]. However, *Gdf15* mRNA levels increased readily with induction of metabolic acidosis in the CNT, CCD and mostly OMCD. After 3 days of acid load, *Gdf15* mRNA was maximally increased and declined with prolonged acidosis. Strikingly, the time-course of *Gdf15* mRNA regulation upon acid loading matched the relative proliferation rates of type A-ICs. Moreover, acidotic Gdf15 deficient mice showed a reduced and delayed proliferation rate of type A-ICs [146].

Interestingly, the early proliferation of type A-ICs of acidotic mice was shown to be dependent on the regulation of the cell-cycle regulatory protein Cyclin D1, which is a downstream target of the phosphatidylinositol-3 kinase (PI3K)/Akt/mammalian target of rapamycin (mTOR) signalling pathway. Consistently, the cell-cycle regulatory

proteins p53 and cyclin D1/D3 were abnormally regulated in Gdf15 KO mice. The molecular action of Gdf15 in renal CD differentiation remains uncertain. However, a putative role in controlling the orientation of the mitotic spindle by Gdf15 is suggested. Hence, this study [146] demonstrated nicely, that metabolic acidosis induces the proliferation of terminally differentiated type A-IC population in the renal OMCD of mice. Notably, proliferation markers were rarely detected in PCs or type B-ICs. In addition, the data indicate that p53, Gdf15 and the PI3K/Akt/mTOR regulatory pathway are required for the acute proliferation event. However, detailed investigation of the interactive mechanisms is needed.



**Fig. 5.2 Illustration of cellular processing, secretion and storage of Mic-1 (Gdf15)**, taken from [181]. The biologically active mature Mic-1 protein is a secreted dimer and induces distinct autocrine and paracrine actions upon various stimuli.

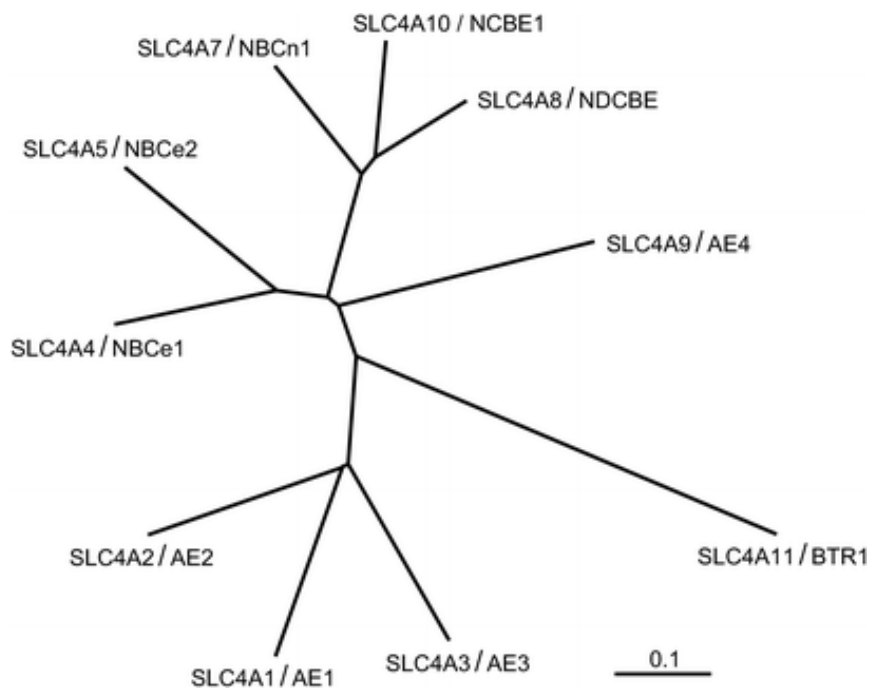
## 6. BICARBONATE TRANSPORTER

At present, two evolutionary unrelated gene families encoding for bicarbonate transporting proteins were identified in humans, the SLC4 (solute carrier 4) family of anion exchangers (AE) and the SLC26 family of sulfate permease anion transporters [182]. The gene products of either family display distinct phylogenetic relationships with differences in their tissue and subcellular distribution, as well as substrate preference, transport mode and regulatory mechanisms. Deficiencies in some of the members of both families lead to hereditary diseases affecting different organ functions and thus display characteristic phenotypes [127].

### 6.1 *The SLC4 Family*

In humans, the SLC4 (solute carrier 4) family of  $\text{Cl}^-/\text{HCO}_3^-$  exchangers consists of ten members with distinct phylogenetic relationships. They are distinguished by their transport mechanism and preferred substrate and thus belong to at least three different major clades (Fig. 6.1) [183]. The  $\text{Na}^+$ -independent and electroneutral  $\text{Cl}^-/\text{HCO}_3^-$  exchangers SLC4A1 (AE1), SLC4A2 (AE2), and SLC4A3 (AE3). The  $\text{Na}^+$ -dependent electrogenic  $\text{HCO}_3^-$  cotransporter SLC4A4 (NBCe1) and SLC4A5 (NBCe2), and the  $\text{Na}^+$ -dependent electroneutral  $\text{HCO}_3^-$  cotransporter SLC4A7 (NBCn1) and SLC4A10 (NBCn2), as well as the  $\text{Na}^+/\text{HCO}_3^-/\text{Cl}^-$  exchanger SLC4A8 (NDCBE). SLC4A9 (AE4) does also belong to the  $\text{Na}^+$  dependent clade but its transport function and substrate preference are still unresolved. SLC4A11 (BTR1) belongs to the third major clade, since it shares similarities with borate transporters of plants, and fungi. However, its transport mechanism and anion preferences remain elusive.

SLC4 homologues were identified in genomes of many different organisms, such as marine bacteria, plants, fungi, marine invertebrates, insects, and a variety of vertebrate and mammalian species [183].



**Fig. 6.1 Phylogenetic tree of human SLC4 family**, taken from [183].

Members of the SLC4 family are crucial for several physiologic processes in mammals, such as the carriage of  $\text{CO}_2$  from tissue to lungs, epithelial secretion and reabsorption of acid-base equivalents (e.g. in the kidney, stomach and pancreas), reabsorption of NaCl (e.g. ileum and proximal colon) as well as regulation of cell volume and intracellular pH in almost every cell of the body [183].

All SLC4 polypeptides share three common structural domains and features. They have a remarkably long intracellular hydrophilic N-terminal domain consisting of 400-700 amino acids, followed by a polytopic transmembrane domain of ~500 amino acids, which spans the membrane 10-14 times, and a short cytoplasmic C-terminal domain of 30-100 amino acids [184]. The predicted topology and structure of SLC4 transporters is mostly derived from studies of the highly abundant human erythroid AE1 protein (Fig. 6.2). Almost all *SLC4* genes express 5'-variant transcripts from alternate promoters, thus generating different polypeptide isoforms with distinct N-terminal domains.

### **6.1.1 Electroneutral Anion Exchangers**

Among the SLC4 family, three members (AE1, AE2 and AE3) belong to the clade of electroneutral and  $\text{Na}^+$ -independent  $\text{Cl}^-/\text{HCO}_3^-$  exchangers. As they mediate the 1:1 exchange of monovalent anions from opposing sites of the plasma membrane, they are also referred to as anion exchangers (AEs). Their preferred substrates are  $\text{Cl}^-$  and  $\text{HCO}_3^-$ , but they can also transport  $\text{OH}^-$ . In addition, AE1 was shown to mediate the exchange of  $\text{SO}_4^{2-}$  and  $\text{H}^+$  for  $\text{Cl}^-$  at very low rates [185]. Since the anion exchange mechanism is driven by substrate gradients, AEs can mediate influx and efflux of  $\text{HCO}_3^-$  by reversing the transport mode, but only erythroidal AE1 is known to work reversibly under physiological conditions [186]. Pharmacologically, all AEs are inhibited by the disulfonic stilbenes SITS and DIDS. However, the oxonol dye diBA5(C4) was shown to be a more potent inhibitor of AE1 than AE2, whereas polyaminosterols were found to inhibit AE2 more potently than AE1 [187].

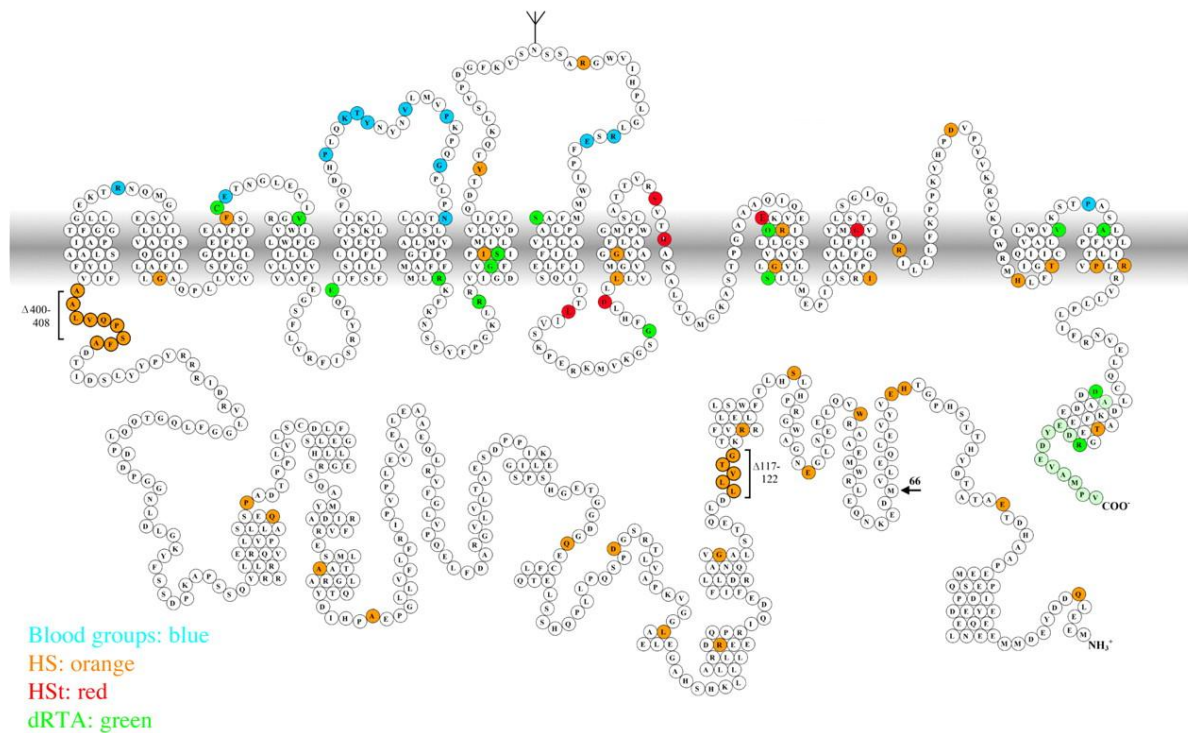
The structure of the three AEs is highly conserved, especially within the transmembrane (TM) domains, as they share up to 56% of amino acid identity [183]. However, they are distinct in their regulation and transport activity. Interestingly, in a physiological range, AE1 activity is not modulated by pH, whereas AE2 and AE3 activity is highly sensitive to changes in pH [182, 188]. Additionally, AE2, but not AE1, was shown to be activated by hypertonicity and contributes to cell volume regulation when expressed in *Xenopus laevis* oocytes [182].

Furthermore, AE1 has a single N-glycosylation site, of yet unknown function, at the fourth extracellular loop, while AE2 and AE3 are N-glycosylated at the third extracellular TM domain [184]. However, the N-glycosylation of AE2 and AE3 had only little effects on transport activity or membrane expression [189].

The present study focuses on the relevance of Ae1 in the kidney. Notably, renal deficiency of Ae1 does not seem to induce compensatory regulation of other AEs in mouse kidney [190].

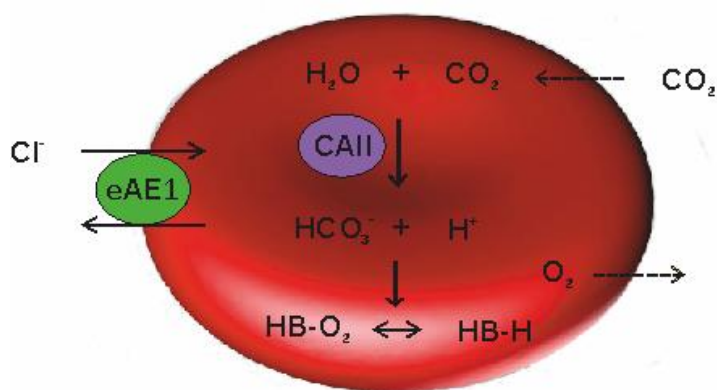
### **6.1.2 The Anion exchanger 1**

In mammals, the anion exchanger 1 (AE1) is predominantly expressed in the plasma membrane of red blood cells (RBCs) and the basolateral membrane of renal type A-ICs (Fig. 3.3). Lower expression levels of AE1 are also present in the heart, distal colon, glomerular podocytes and other tissues [191, 192]. Homologous of mammalian AE1 were also identified in insects and amphibians [51, 193, 194]. The *AE1* gene encodes two polypeptide variants, the longer erythroid AE1 (eAE1, red cell band 3) and the shorter kidney AE1 (kAE1). In humans, kAE1 initiates at Met66 (methionine 66), while it initiates at Met79 in mice [184]. Numerous mutations of the *SLC4A1* gene have been identified in humans, but also cattle and zebrafish, which either affect the structure and function of RBCs or the kidney (Fig. 6.2) [184].



**Fig. 6.2 Predicted topology model for human AE1 polypeptide**, after Zhu et al. [195] taken from [184]. Polymorphisms encoding blood group antigens are highlighted in blue. Mutations (missense, nonsense, splicing and deletion) associated with hereditary spherocytic anemia and ovalocytosis (HS) are highlighted in orange, missense mutations leading to hereditary stomatocytosis (Hst) and xerocytosis are shown in red. Mutations identified in dominant and recessive distal renal tubular acidosis (dRTA) are labeled in green and terminal truncations are in lighter orange and green.

In the RBC, eAE1 is the most abundant integral membrane protein and serves, together with the action of the metalloenzyme carbonic anhydrase II (CAII), to increase the CO<sub>2</sub> carriage capacity of the blood and thus its removal by the lungs (Fig. 6.3). CAII was suggested to interact with the short C-terminus of eAE1 forming a pH dependent functional transport metabolon that accelerates transmembrane bicarbonate transport [196, 197].

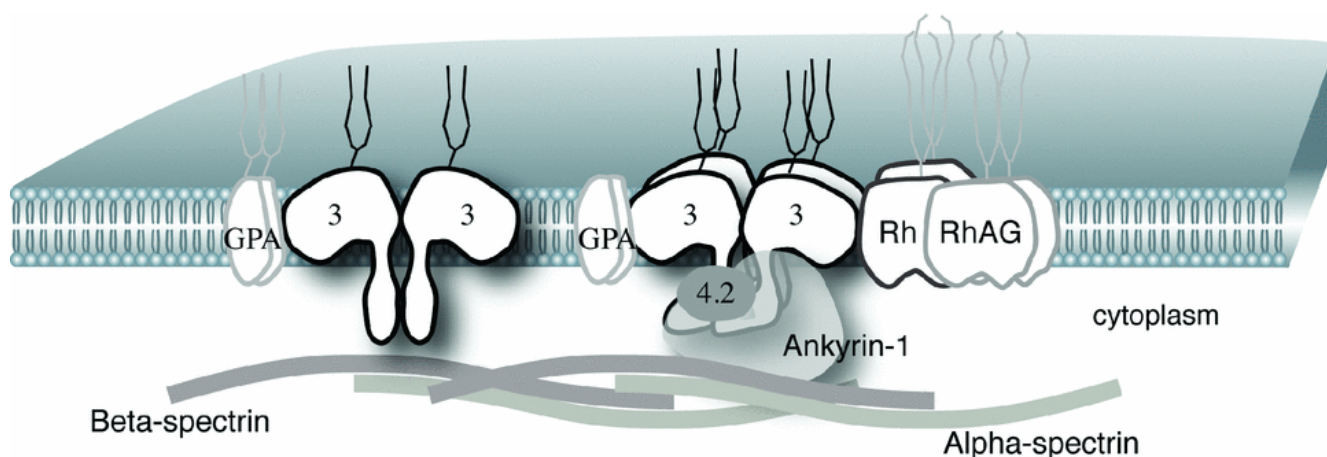


**Fig. 6.3 Role of AE1 in erythroid CO<sub>2</sub> carriage (Jacobs-Stewart Cycle),** modified from [183]. In the capillaries, CO<sub>2</sub> enters the erythrocyte mostly by transmembrane diffusion and probably via the water channel aquaporin 1. Soluble CA II (carbonic anhydrase II) catalyzes the rapid formation of HCO<sub>3</sub><sup>-</sup> from intracellular CO<sub>2</sub> and H<sub>2</sub>O. Deoxygenated hemoglobin buffers excess H<sup>+</sup> (Hb-H), while eAE1 releases newly generated HCO<sub>3</sub><sup>-</sup> into the blood plasma in the exchange for extracellular Cl<sup>-</sup> ("Hamburger"-Shift). The cycle is reversed in the lungs with oxygenation of hemoglobin (Hb-O<sub>2</sub>).

Furthermore, the presence of eAE1 dimers or tetramers in the RBC membrane is known to be crucial for the stability and integrity of the membrane as it is a central part of a cytoskeletal macrocomplex (Fig. 6.4) [186]. In the RBC, the N-terminal domain of eAE1 anchors the spectrin based cytoskeleton to the plasma membrane via interaction with ankyrin. Moreover, eAE1 was shown to interact with protein 4.1 and 4.2, hemoglobin, glycolytic enzymes (aldolase, GAPDH, protein tyrosine kinase) and rhesus glycoprotein complexes [198, 199].

However, possible interactions of kAE1 with other proteins in the renal type A-ICs are less well understood and most likely different, since the kAE1 N-terminal domain lacks the first 65 amino acids in humans or 78 amino acids in mouse and therefore the ability of, for example ankyrin binding. Additionally, mutations of AE1 affecting both organs simultaneously, RBCs and kidney, are rarely seen.





**Fig. 6.4 AE1 macrocomplex in the erythroidal membrane**, taken from [186]. In the RBC membrane, eAE1 (indicated as band 3) is present as dimers or tetramers, which interact with other cytoskeletal proteins forming a macrocomplex. Dimers of eAE1 bind to glycophorin A (GPA), while tetrameric eAE1 is further interacting with the rhesus glycoprotein RhAG and the cytoskeletal complex protein 4.2, ankyrin-1 and  $\alpha$ - and  $\beta$ - spectrin.

Several mutations of human AE1 are identified and indicated in Fig. 6.2. They give rise to erythrocyte malformation and malfunction with haemolytic anaemia in dominant hereditary spherocytosis (HS) or stomatocytosis (Hst) with overhydrated cation-leak syndrome, or affect the kidney function leading to primary dRTA [200]. Notably, mutations leading to HS are distributed throughout the whole transporter, while Hst mutations are restricted to the first part of the second half of the TM domain and dRTA associated mutations are spread within the TM domain and the C-terminal tail. Since the *SLC4A1* gene generates at least two tissue specific products, mutations affecting both organs simultaneously are rarely seen. However, dominantly inherited mutations affecting the RBCs might be recessively inherited for renal defects [186] and dominantly inherited RBC mutants are associated with 50% expression of the wildtype allele, which seems to be sufficient for normal renal function [200]. Additionally, some dRTA associated AE1 mutations are rescued by chaperones like glycophorin A (GPA), which are expressed in RBCs or heterologous expression systems, but not in kidney [201-203].

In the kidney, kAE1 is abundantly expressed at the basolateral membrane of type A-ICs (Fig. 2.6, 3.2, 3.3). Together with apical  $H^+$  secretion, kAE1 is required for the acidification of the final urine, as it is the major basolateral bicarbonate extruder. Deficiency for kAE1 in man, mouse and cattle leads to dominant or recessive forms of dRTA. Among humans, mutations such as kAE1 R589 (H, C, S), S613F and R901X were identified in dominant dRTA, while G701D, S773P and S667F cause recessive forms. Several *in vitro* studies of dRTA associated AE1 mutants try to explain their pathogenicity and give rise to possible mechanisms involved in renal AE1 biosynthesis and targeting. Interestingly, the dominant negative kAE1 R901X mutation was shown to be mistargeted to the apical membrane or retained intracellularly in MDCK cells [204, 205]. Similarly, expression of dominant negative kAE1 R589H and S613F in polarized MDCK cells showed the retention of the protein within the ER (endoplasmic reticulum) as well, where it can dimerize with the wild-type kAE1 and thus prevent its trafficking to the surface [206]. However, when expressed in *Xenopus* oocytes, kAE1 R589H retains partial activity and surface expression [207], which might be due to incubation with lower temperature (temperature-sensitive folding mutation), or the presence of different trafficking, signalling and folding chaperones in *Xenopus* derived cells [200]. Anyhow, in a renal biopsy from a patient with dominant kAE1 R589H associated dRTA and superimposed pyelonephritis and nephrocalcinosis, a reduced number of intercalated cells was observed, with preserved expression of V-ATPase, but absence of detectable kAE1 [208]. In contrast, analysis of a renal biopsy from a patient with the dominant kAE1 S613F mutation, showed reduced number and size of ICs, with an exclusively cytosolic expression of kAE1 [138].

## **6.2 The SLC26 Family**

The SLC26 family of anion exchangers consists of ten members that play critical roles in a variety of physiological processes, such as skeletal development, synthesis of thyroid hormone, transepithelial  $\text{Na}^+\text{-Cl}^-$  transport, renal bicarbonate excretion and pancreatic bicarbonate secretion [209]. Accordingly, SLC26 transporters are very distinct in their expression pattern. Moreover, they transport a variety of monovalent and divalent anions, including sulfate, chloride, iodide, formate, oxalate, hydroxyl ion, and bicarbonate with variable transport specificity [209].

Within the kidney, several SLC26 members are expressed along the renal tubule (i.e. SLC26A1, SLC26A4, SLC26A7) [209-211]. However, the present study focuses exclusively on the  $\text{Cl}^-/\text{HCO}_3^-$  exchanger Pendrin (SLC26A4).

### **6.2.1 Pendrin**

Pendrin (PDS, SLC26A4) is expressed in the inner ear, thyroid gland, and the kidney and transports chloride, bicarbonate, and iodide [33, 209]. In humans, mutations of PDS are associated with Pendred Syndrome, which is clinically characterized by hypothyroidism and goiter, as well as sensorineural deafness [33]. In the kidney, Pds is exclusively expressed on the luminal membrane of bicarbonate secreting type B-ICs and type non-A/non-B ICs of the CD system (chapter 3.4, Fig. 2.6 and 3.4). As described in chapter 2.5, Pds expressing cells are restricted to the renal cortex of adult kidneys (Fig. 2.4, 2.5). However, during renal development Pds expressing cells derive from both, the MM and UB pool of progenitor cells (Fig. 2.5).

The expression of Pds in mouse kidney and inner ear was shown to depend on the transcriptional activation by the forkhead box protein Foxi1 (chapter 2.7.3). In adult rodent kidneys Pds expression is highly regulated by systemic acid-base status (chapter 4.4, 5.2, and 5.2.1) and systemic chloride levels (for review see [33]). However, humans and mice with genetic disruption of Pds have normal acid-base

balance under basal conditions [33, 212]. Nevertheless, two independent Pds deficient mouse models show a reduced urinary pH due to a reduced urinary buffering of  $H^+$  by secreted  $HCO_3^-$  [212, 213]. Interestingly, the abundance of type non-A ICs, but not A-ICs, was shown to be reduced in the renal cortex of Pds KO mice [212]. Moreover, type B-ICs in Pds deficient kidneys were markedly reduced in size and their ultrastructural appearance was altered [212]. Concomitantly, a reduction of transporters involved in renal  $H^+/OH^-$  handling (i.e. V-ATPase, NBC3, Rhbg) was noted, probably attenuating a rise in intracellular and systemic pH due to the loss of Pds [212].

## **7. MOUSE MODELS USED IN THIS STUDY**

In the present study, distinct mouse models were used to answer the question whether an acid-base disturbance *per se* or the absence of renal Ae1 would introduce the observed abnormal presence of IC marker in Ae1 deficient mice.

### **7.1 *Mouse models with primary dRTA***

As already described in detail, renal Ae1 deficiency leads to distal renal tubular acidosis (dRTA) in mouse and men. Furthermore, loss of function mutations or absence of Atp6v1b1 and Rhcg in mice was also shown to reduce the kidneys ability of maximal urinary acidification and therefore the removal of systemic excess acid. Hence, a genetic defect in one of the three proteins impairs renal IC mediated acid-base handling and thus causes primary dRTA. Therefore, Atp6v1b1 and Rhcg deficient mice serve as alternative models to investigate a possible impact of inherited dRTA on the differentiation of ICs, respectively.

#### **7.1.1 *Atp6v1b1 deficient mice***

Deficiency of the B1 subunit of the V-ATPase (ATP6V1B1) leads to spontaneous dRTA in humans [214]. Nevertheless, the phenotype is different in mice, as a homozygous deficiency of the B1 subunit of the V-ATPase (Atp6v1b1) leads to incomplete dRTA [215]. Briefly, under standardized laboratory conditions and feeding with normal rodent chow B1 KO mice did not show a detectable renal acid-base disturbance. Consistently an increased abundance of the alternative B subunit (B2) of V-ATPase was detected, which might partially compensate for B1 deficiency. Anyhow, after an oral acid load these animals develop systemic acidosis, due to an insufficient urinary acidification. It seems reasonable, that the strong alkali load provided by the standard rodent chow, might help to compensate insufficient urinary acidification under basal conditions [215].

### **7.1.2 *Rhcg* deficient mice**

Distal urinary acidification is a complex process, involving the coordinated secretion of protons by luminal V-ATPase and the recovery of bicarbonate to the blood by basolateral  $\text{Cl}^-/\text{HCO}_3^-$  exchanger. As already discussed, this process is limited and thus additional alternate acid extrusion pathways like titratable acids and essentially ammonium are required. Recently, the rhesus glycoprotein Rhcg was identified as an ammonia transporter in mice, rats and humans [109]. Similar to murine B1 deficiency, mice lacking Rhcg display incomplete dRTA, as unravelled by oral acid load test [114]. Again, animals under basal conditions were raised with standard rodent chow, implicating large alkali loads that may compensate for renal insufficiency. Two independently generated Rhcg KO mice are available (Marini AM, Bruxelles [114] and TIGM, USA (Bourgeois, Bounoure, Christensen, Devuyst, Wagner, unpublished data)). However, both mouse models develop incomplete dRTA with minor differences under basal conditions.

## **7.2 *Induced chronic metabolic acidosis model***

As reported above, B1 and Rhcg deficient mice are prone to develop dRTA under forced acid-base disturbances. However, Ae1 KO mice and patients with dominant mutations of kAE1 suffer chronically from complete dRTA. Experimentally induced acute and particularly chronic acid-base disturbances in rodents were observed to alter the renal CD morphology (chapter 5.2, 5.2.1 5.2.2). Thus, the observed dysregulation of CD cell markers in Ae1 deficient subjects might be a result from the chronic acid-base disturbance or from the loss of Ae1 protein in the kidney. The oral application of  $\text{NH}_4\text{Cl}$  to the food or drinking water of rodents is a common experimental approach to induce metabolic acidosis [155, 216, 217]. Even though a more than 7 days lasting treatment with  $\text{NH}_4\text{Cl}$  is already considered a chronic acid-base disturbance and sufficient to cause adaptive responses in renal CD morphology, the period of time does not compare with the lifelong deteriorating dRTA in Ae1 deficient subjects. Furthermore, an altered differentiation of CD markers was initially observed in adult (P84) Ae1 KO mice. Therefore, metabolic acidosis in C57/Bl6 wildtype mice was induced for a total of 8 weeks, starting right away after weaning at 3 to 4 weeks of life. Based on recent studies, the supplementation of equivalent amounts of  $\text{NH}_4\text{Cl}$  to the food was favoured over the addition of 0.28M  $\text{NH}_4\text{Cl}$  to the drinking water to avoid unwanted side effects by chronic dehydration of the animals [218, 219].

## **8. AIM OF THE STUDY**

Similar to humans, Ae1 deficiency in mice is correlated with the development of complete dRTA and nephrocalcinosis, as well as dysmorphology and malfunctionality of erythrocytes [190, 200, 220]. Furthermore, a severe urinary concentrating defect associated with dysregulated renal expression and localization of the water channel Aqp2 was observed in Ae1 deficient mice [190]. Additionally, unpublished observations indicated an abnormal distribution of intercalated cell marker proteins within the renal CD system of Ae1 deficient mice. Moreover, in renal biopsies of patients carrying dRTA associated AE1 mutations (AE1R589H and AE1S613F) aberrancies of V-ATPase expressing cells in the renal CDs have been reported [138, 208].

Therefore, the major aim was to examine the role of Ae1 in collecting duct cell differentiation by addressing the following subaims:

- 1) Characterization and quantification of possible abnormal expression pattern of renal IC marker by immunofluorescence (IF) in Ae1 deficient mice at different developmental and mature time points of life.
- 2) Investigation of a possible impact of acid-base disturbance on IC differentiation, by comparing the abnormal IC marker expression in Ae1 deficient mice with expression profiles in distinct mouse models for incomplete dRTA and chronically induced metabolic acidosis (chapter 5.2.1).
- 3) Immunologic comparison of IC marker expression in Ae1 deficient mice and patients carrying AE1 mutations associated with dRTA.
- 4) Investigation of the possible impact of candidate molecules (transcription factors, signalling molecules, receptors), which have been reported to play a role in IC development and differentiation (chapter 2.7 to 2.7.6, 5.2.2), on the abnormal IC differentiation observed in Ae1 deficient mice, by IF, qRT-PCR and immunoblot.



## 9. MATERIAL AND METHODS

### 9.1 *Animal experiments*

Animal experiments were performed in the ZIHP (Zurich Center for Integrative Human Physiology) Core facility for Rodent physiology (ZIRP) according to the rules and guidelines of the Swiss animal welfare law and approved by the Veterinäramt Kanton Zürich, Switzerland.

Ae1 deficient mice were obtained from Luanne Petersen and Seth Alper, Boston, Massachusetts, USA [220]. B1 deficient mice were obtained from Richard P. Lifton, New Haven, Connecticut, USA [215]. Rhcg deficient mice were obtained from TIGM (Texas A&M Institute for Genomic Medicine). All transgenic strains were bred in heterozygous pairs in the in-house animal facility of the University of Zurich.

Where appropriate, time of gestation was assessed by monitoring vaginal mating plugs. Adult mice or embryos and pups were euthanized by cervical dislocation or decapitation for organ collection. Both kidneys were subsequently removed, freed from the renal capsule, cut into halves, snap frozen in liquid nitrogen (N<sub>2</sub>) and stored at -80°C until further analysis.

C57/Bl6 male mice were obtained from Institute of Veterinary Sciences, Fuellinsdorf, Switzerland at the age of 3 to 4 weeks and exposed to an oral acid load for 8 weeks to induce chronic metabolic acidosis. The animals were housed in normal cages with tap water *ad libitum* and fed standard rodent powder chow (GLP 3433, Kliba Nafag, Switzerland). Supplementation of 2 g NH<sub>4</sub>Cl/100 g powdered food (equivalent to 0.28 M NH<sub>4</sub>Cl in the drinking water) for 8 weeks was given to induce chronic metabolic acidosis [170, 219]. The bodyweight of all animals was assessed weekly and the health status (indicated by the appearance of the fur and activity) was determined at least every 2 days. At the end of the experimental period, the animals were placed in metabolic cages (Tecniplast, Italy). The mice were allowed to adapt to

the new environment for 3 days before assessing metabolic parameters and collection of 24 hours urine samples under mineral oil for another 2 days.

Finally, mice were anesthetized with 1.5% Isoflurane/pressurized air and 500  $\mu$ l venous blood was taken with a heparinised syringe from the *V. cava caudalis* for subsequent blood gas analysis (ABL505, Radiometer, Copenhagen). The remaining blood was centrifuged at 5,000 rpm (4°C) for 5 minutes and the obtained plasma was snap frozen in liquid N<sub>2</sub> for further analysis. Finishing blood sampling, 5 animals per group were perfused with phosphate buffered saline (PBS, pH 7.4) through the left ventricle to allow wash-out of residual blood. Both kidneys were subsequently removed, freed from the capsule, cut into halves and snap frozen in liquid N<sub>2</sub> for further analysis. Another 5 animals per group were processed for organ fixation as described in 9.2.

## **9.2 Fixation and cryopreservation of whole organs**

Adult and juvenile mice were anesthetized with 1.5 % Isoflurane/pressurized air and perfused through the left ventricle with 10 ml preperfusate consisting of 10,000 IU Heparin-Na Solution (BBraun), 2 ml Rapidocain 1 % (Ratiopharm), 2 ml CaCl<sub>2</sub> 16 %, 2 ml 0.9 % NaCl and 2 ml Aqua destillata. Subsequently, the animal was perfusion fixed with 3 % Paraformaldehyde (PFA) in 0.1 M Na-cacodylate buffer. After 5 minutes the kidneys were removed and postfixed for an additional hour on ice in the above fixative solution. Embryos and pups were euthanized by decapitation and kidneys were subsequently removed and immersion fixed in 3 % PFA/0.1 M Na-cacodylate buffer for 1 hour at 4°C. Following fixation, the kidneys were stored at 4°C in 1% PFA/0.1 M Na-cacodylate buffer. Cryoprotection was assured by immersion in 30% Sucrose/PBS. The kidneys were then embedded in Tissue-Tek Cryomolds (VWR) and Neg-50 frozen section medium (Richard-Allan Scientifics) with liquid propane and stored at -80°C.

### **9.3 Immunofluorescence**

Paraffin embedded human kidney biopsy sections (kind gifts of Robert Unwin, University College London, UK and Seth Alper, Beth Israel Deaconess Medical Center, Boston, MA) were dewaxed in a series of 2 x 10 min 100 % Xylol, 3 x 3 min 100 % ethanol (EtOH), 3 x 3 min 96 % EtOH, and 1 x 30 min 75 % EtOH before rehydration in PBS. Cryopreserved mouse kidneys were cut into 3-5  $\mu$ m thin slices (Cryostat CM1850, Leica) and rehydrated in PBS for at least 60 minutes. Different antigen retrievals were carried out, where necessary. Paraffinized tissue sections were treated by microwaving for 4 minutes in 10 mM citric acid pH 6.0 and 1 % sodium dodecyl sulfate (SDS) in PBS for 5 minutes. Cryopreserved tissue sections were either treated with 1 % SDS/PBS for 5 minutes, 0.3 % Triton X/PBS for 15 minutes or by microwaving. Following antigen retrieval, the sections were washed in PBS for 3 x 5 minutes. Unspecific binding of antibodies was impeded by 15 minutes incubation with 1 % bovine serum albumin (BSA) in PBS. Antibodies were diluted in 0.02 % Na-azide/PBS and stored at 4°C. Primary antibodies were incubated for overnight at 4°C in a moist chamber followed by two washes with hypertonic PBS (18 g/l NaCl/PBS) and 1 x wash with PBS. Secondary antibodies containing 4',6-diamidino-2-phenylindole (DAPI) were incubated in the dark for 120 minutes at room temperature (RT) in a moist chamber followed by previously described washing-steps. The stained sections were finally embedded with Glycergel (DakoCytomation) and kept at 4°C in the dark until microscopic analysis. Photomicrographs were taken using an upright DM 5500 fluorescence microscope with AF 6000 System (Leica) including the objectives HC PL FLUOTAR 10.0x0.30 DRY, HCX PL APO 40.0X1.25 OIL and HCX PL FUOTAR 100.0X1.30 OIL. Fluorescent specimens were pictured using a DFC 360 FX monochrome camera (Leica) and chromogenic specimens were pictured by bright-field microscopy using the digital colour camera DFC 295 (Leica). Images were further analyzed and processed using the LAS AF Lite Software (Leica), Corel PHOTO-PAINT X5 and the open-source software Image J (National Institute of Health).

The following combinations of primary antibodies detecting different marker proteins were used in distinct colocalization studies to distinguish principal cells, variations of ICs, type A-ICs and type B-ICs (Tab. 9.1).

**Tab. 9.1**      **Marker combinations in immunofluorescent colocalization studies**

<b>Species: Mouse</b>			
1)	Aqp2	Pds	Calb1
2)	Aqp2	Ae1	Calb1
3)	Aqp2	a4	Calb1
4)	Aqp2	B1	Calb1
5)	Aqp2	CaII	Calb1

<b>Species: Human</b>			
1)	AQP2	PDS	AE1
2)	AQP2	PDS	a4
3)	AQP2	PDS	B1
4)	AQP2	PDS	CAII

### **9.5    *Quantification of cells***

Immunofluorescently labelled cells from at least one kidney cross section per animal and each set of antibody combination was counted (Tab. 9.1). Principal cells were identified by luminal aquaporin 2 expression. Intercalated cells were either labelled for the a4 or B1 subunit of the V-ATPase. Type B-ICs were identified by the apical presence of Pds and type A-ICs by the basolateral presence of Ae1, where appropriate. The cellular affiliation to distinct renal tubule segments was taken into account. The renal CNT, CCD, OMCD and IMCD were primarily distinguished based on morphologic criteria. Immunologic presence of cytosolic calbindin D28K (Calb1) was further used to discriminate between the CNT and CCD segment [21].

**Tab. 9.2 List of Antibodies (AB) used for immunofluorescence**

<b>1° AB</b>	<b>Antigen/Catalog No.</b>	<b>dilution (mouse)</b>	<b>dilution (human)</b>	<b>Source</b>
rabbit-anti-mouse Ae1	DCFIYEDQIRPQDREELLRA LLK	1:1,000	1:500	Own (Pineda)
guinea pig-anti-mouse Ae1	DCFIYEDQIRPQDREELLRA LLK	1:500		Own (Pineda)
rabbit-anti-mouse Pendrin	CKDPLDLMEAEMNAEELD VQDEAMRRLAS	1:1,000		Own (Pineda)
guinea pig-anti-mouse Pendrin	CKDPLDLMEAEMNAEELD VQDEAMRRLAS	1:1,000	1:250	Own (Pineda)
rabbit-anti-a4	CKFSPFSFKHILDGTAE	1:1,500	1:500	Own (Pineda)
rabbit-anti-B1	CAQQDPASDTAL	1:1,000	1:250	Own (Pineda)
guinea pig-anti-B1	CAQQDPASDTAL	1:500		Own (Pineda)
rabbit-anti-CaII	AB1828	1:1,500	1:500	Chemicon Int.
goat-anti-Aqp2	sc9882	1:2,000	1:200	Santa Cruz
mouse-anti-Calbindin D28K	McAB 300	1:20,000		Swant
goat-anti-Foxi1	TSGVLYPREGTEV	1:200		abcam
rabbit-anti-Cp211	MLFWHTQPEHYNQHNSG SC	1:1,500		[90]

<b>2° AB</b>	<b>Antigen/Catalog No.</b>	<b>dilution (mouse)</b>	<b>dilution (human)</b>	<b>Source</b>
donkey-anti-goat IgG Alexa 647	A21447	1:1,000	1:500	Invitrogen
donkey-anti-goat IgG Alexa 488	A11055	1:1,000		Invitrogen
donkey-anti-rabbit IgG Alexa488	A21206	1:1,000	1:1,000	Invitrogen
donkey-anti-rabbit IgG Alexa594	A21207	1:1,000		Invitrogen
donkey-anti-guinea pig IgG Cy2	706-226-148	1:500		dianova
donkey-anti-guinea pig IgG Cy3	706-165-148	1:500	1:500	dianova
donkey-anti-guinea pig IgG Dylight 649	706-496-148	1:500		Jackson Immuno Lab
donkey-anti-mouse IgG Dylight 649	715-496-151	1:500		Jackson Immuno Lab
donkey-anti-mouse IgG Alexa594	A21203	1:1,000		Invitrogen
donkey-anti-mouse IgG Alexa488	A21202	1:1,000		Invitrogen
4',6-diamidino-2-phenylindole, dilactate (DAPI, dilactate)	D3571	1:1,000	1:1,000	Invitrogen

## 9.6 RNA Isolation and qRT-PCR

Total RNA was isolated from half a kidney of adult mice (P30 and P84) using the RNEasy mini Kit (Qiagen). Total RNA from one kidney of mouse embryos (E18) or newborn mice (P3) was isolated using the RNEasy micro Kit (Qiagen). RNA isolation and on-column DNA digestion was performed according to the manufacturer's protocols. Total RNA quantity was measured using a ND-1000 UV/Vis-Spectrophotometer (NanoDrop Technologies). Reverse transcription of total RNA (300 ng) was carried out using the TaqMan Reverse Transcription Kit (Applied Biosystems). Quantitative real-time PCR (qRT-PCR) was performed with an ABI PRISM 7700 sequence detection system (Applied Biosystems) using TaqMan Universal PCR Master Mix (Applied Biosystems). Specific primers for the genes of interest were designed using the online-tool Primer3 (<http://frodo.wi.mit.edu/primer3/>) and tested by conventional PCR to generate the expected amplification products. Oligonucleotide probes conjugated to the reporter dye FAM at the 5' end and the quencher TAMRA at the 3' end were obtained from Microsynth, Switzerland. The mRNA expression levels of genes of interest were firstly normalized to the expression levels of the housekeeping gene *hypoxanthine-guanine-phosphoribosyltransferase* (*Hprt*). Secondly, the normalized values were calculated in percentage and statistical analysis was performed, respectively.

Reverse Transcription			Real Time RT-PCR			
Step	Temperature	time	Step	Temperature	time	cycle
1	25°C	10 min	1	50°C	2 min	
2	40°C	30 min	2	95°C	10 min	
3	95°C	5 min	3	95°C	15 sec	40
4	4°C	hold	4	60°C	1 min	
			5	4°C	hold	

**Tab. 9.3 List of primer and Taqman probes used for qRT-PCR**

<b>Gene</b>	<b>Accession No.</b>	<b>forward primer</b>
<i>Hprt</i>	NM_013556	5'- TTATCAGACTGAAGAGCTACTGTAATGATC -3'
<i>Aqp2</i>	NM_009699	5'- TGGTGCTGTGCATCTTTGCCT -3'
<i>Slc4a1 (Ae1)</i>	NM_011403	5'- AGGACCTGGTGTGCCAGAG -3'
<i>Slc26a4 (Pds)</i>	NM_011867	5'- GCCTTTGGGATAAGCAACGTC -3'
<i>Cp2l1</i>	NM_023755	5'- ATGCTGTTCTGGCACACGCAGC -3'
<i>Foxi1</i>	NM_023907	5'- AAGAAGGTGCCCCGAGATG -3'
<i>Klf4</i>	NM_010637	5'- CTGCCAGACCAGATGCAGT -3'
<i>Bdkrb2</i>	NM_009747	5'- ATGCCCTGCTCCTGGAAGCTACT -3'
<i>Gdf15</i>	NM_011819	5'- AGAGGACTCGAACTCAGAACCAAG -3'
<i>Rhcg</i>	NM_019799	5'- GTTGAGAAGAAGCGCAAGAA -3'
<i>Atp6v1b1 (B1)</i>	NM_134157	5'- AGGACAGTGTGCAGCGTCAAT -3'
<i>Hensin (Dmbt1)</i>	NM_007769.2	5'- GCTCTTTCACCTCAACATCCA -3'

<b>Gene</b>	<b>Accession No.</b>	<b>reverse primer</b>
<i>Hprt</i>	NM_013556	5'- TTACCAGTGTCAATTATATCTTCAACAATC -3'
<i>Aqp2</i>	NM_009699	5'- ACTTGCCAGTGACAACTGCTG -3'
<i>Slc4a1 (Ae1)</i>	NM_011403	5'- CGGTTATGCGCCATGGA -3'
<i>Slc26a4 (Pds)</i>	NM_011867	5'- CAACGATGGCAACATCACA -3'
<i>Cp2l1</i>	NM_023755	5'- TCTCAGGAGATAGCTGCGGCTC -3'
<i>Foxi1</i>	NM_023907	5'- TCTTCTCACAGTTAGGGTCCAGAGT -3'
<i>Klf4</i>	NM_010637	5'- TGGTATAGGTTTTGCCACAGC -3'
<i>Bdkrb2</i>	NM_009747	5'- AGATCTCGGCCACAGTGCAGCTG -3'
<i>Gdf15</i>	NM_011819	5'- TTGACGCGGAGTAGCAGCTGGC -3'
<i>Rhcg</i>	NM_019799	5'- CGAAGACCATGGCGTGTACA -3'
<i>Atp6v1b1 (B1)</i>	NM_134157	5'- CCTGAACAATGGCCTTGGTC -3'
<i>Hensin (Dmbt1)</i>	NM_007769.2	5'- GCTGGCTTGCATGTGATTT -3'

Gene	Accession No.	oligonucleotide (Taqman probe)
<i>Hprt</i>	NM_013556	5'-TGAGAGATCATCTCCACCAATAACTTTTATGTCCC -3'
<i>Aqp2</i>	NM_009699	5'- ACCTCCTTGGGATCTATTTACCCG -3'
<i>Slc4a1 (Ae1)</i>	NM_011403	5'- ACCTGAGGCTCCGCATGTCGACTATACC -3'
<i>Slc26a4 (Pds)</i>	NM_011867	5'- TGGATTTTCTCCTGTTTTGTGGCTACCACT -3'
<i>Cp2l1</i>	NM_023755	5'- TGGCAGCTACTTGCGTGATGTGCTGGC -3'
<i>Foxi1</i>	NM_023907	5'- ACGACCCAGGCAAAGGGAATTACTG -3'
<i>Klf4</i>	NM_010637	5'- TCTCTCCATTATCAAGAGCTCATGCCA -3'
<i>Bdkrb2</i>	NM_009747	5'- GCCTCCTTTGGCATCGAAATGTTC -3'
<i>Gdf15</i>	NM_011819	5'- TGTCCGGATACTCAGTCCAGAGGTGAGA -3'
<i>Rhcg</i>	NM_019799	5'- TTACTATCGCTACCCGAGCTTCCAG -3'
<i>Atp6v1b1 (B1)</i>	NM_134157	5'- CCCAGTATGCTGAGATTGTCAACTTTACCCTCC -3'
<i>Hensin (Dmbt1)</i>	NM_007769.2	5'- TGACAATAACACCACCAATCTCCCTTTGTGAGT -3'

## 9.7 Protein Isolation

Crude membrane and cytosolic proteins were extracted from half a kidney of adult mice (P30 and P84). In detail, the organ was minced on ice and homogenized in 200  $\mu$ l of ice cold resuspension buffer (200 mM Mannitol, 80 mM Hepes, 41 mM KOH, pH 7.5 supplemented with 1 tablet/10 ml Complete Mini Protease Inhibitor Cocktail, Roche) using a Polytron homogenizer (0.5 mm diameter at 20,000 rpm for 1 minute at 4°C). The homogenate was further sonicated on ice before centrifugation at 2,000 rpm for 20 minutes at 4°C. The resulting supernatant was transferred into a prechilled tube and further centrifuged at 41,000 rpm for 1 hour at 4°C. After ultracentrifugation the supernatant (cytosolic proteins) was transferred into a fresh prechilled tube, while the pellet (crude membrane proteins) was resuspended in 50  $\mu$ l of ice cold resuspension buffer. Protein concentration was measured using the Bio-Rad *DC* Protein Assay according to manufacturer recommendations.



In order to obtain total cell lysates, confluent cell monolayers, grown on 6 well plates (Nunclon Surface, nunc), were washed with ice-cold PBS (without  $\text{Ca}^{++}$ ,  $\text{Mg}^{++}$ , Gibco) at 4°C and scraped into a prechilled 1.5 ml Eppendorf tube. The cell suspension was centrifuged at 2,000 rpm for 5 minutes at 4°C and the supernatant was removed. Subsequently, the denssed cell pellet was lysed by addition of 50  $\mu\text{l}$  RIPA buffer (20 mM Tris-HCl pH 7.5, 150 mM NaCl, 1 mM  $\text{Na}_2\text{EDTA}$ , 1 mM EGTA, 1% NP-40, 1% Na-deoxycholate, supplemented with 1 x Protease inhibitor cocktail (Sigma) and 2 mM PMSF) and vigorous pipetting. The crude cell lysate was centrifuged at 10,000 rpm for 10 minutes at 4°C and the supernatant (total cell lysate) was transferred into a fresh tube. Protein concentration was measured using the Bio-Rad Protein Assay following manufacturer recommendations.

## 9.8 Nuclear protein extraction

Buffer A (hypotonic lysis buffer)	
10 mM Tris-Cl, pH 8.0	
150 mM NaCl	
10 mM MgCl <sub>2</sub>	

Homogenization buffer (HB)	
0.27 M Sucrose	in 1x Buffer A
2 mM EDTA, pH 8.0	
0.1 % NP-40	
1 mM PMSF	
0.5 mM DTT	
Protease Inhibitor Cocktail (1 tablet/10 ml, Sigma)	

Nuclear extraction buffer (NEB)	
10 mM Tris-Cl, pH 8.0	
1 mM EDTA, pH 8.0	
400 mM NaCl	
1 mM DTT	
1 mM PMSF	

Sucrose Cushion (SC)	
30 % w/v Sucrose	in 1x Buffer A
2 mM EDTA, pH 8.0	

Nuclear proteins were extracted from half a kidney of adult mice (P30 and P84). The tissue was minced and subsequently homogenized on ice in 2 ml of ice cold homogenization buffer (HB) using a Polytron homogenizer (0.5 mm diameter at 20,000 rpm) for 1 minute. Prior to homogenization, a 3 ml layer of sucrose cushion (SC) was prepared in a 15 ml Sarstedt tube on ice. The homogenate was smoothly transferred to the 15 ml tube overlaying the SC and centrifuged at 2,000 rpm for 15 minutes at 4°C. Subsequently, the supernatant was removed and the denser pellet of nuclei was washed with ice cold PBS and centrifuged again. Following removal of PBS,

the pelleted nuclei were resuspended in 1 volume of ice cold nuclear extraction buffer (NEB) transferred to a prechilled tube and incubated for 15 minutes on ice while vortexing every 5 minutes. Afterwards the samples were centrifuged at 13,200 rpm at 4°C for 15 minutes and the supernatant (nuclear extracts) was transferred into a fresh prechilled tube. Protein concentration was determined using the Bio-Rad Protein Assay according to manufacturer recommendations.

### **9.9 Immunoblotting**

For immunoblotting 20 µg of crude membrane, or cytosolic protein extracts, or total cell lysate, or 5 µg of nuclear protein extracts were solubilized in Laemmli buffer, separated by SDS-PAGE and stained with Coomassie Blue or blotted on a PVDF membrane (Immobilon-P, Millipore). Washing steps were carried out using TBS-T (Tris buffered saline, 0.1 % Tween 20) and unspecific binding of the antibodies was impaired using 5 % non-fat dry milk/TBS-T. Antibodies were diluted according to Tab. 9.4 in 1 % non-fat dry milk/TBS-T. Primary antibodies were incubated overnight at 4°C, while secondary antibodies were incubated for 2 hours at RT. Immunoreactive signals were detected by chemiluminescence (Immobilon Western Substrates, Millipore) using a DIANA III-chemiluminescence detection system (Raytest). Densitometry of the respective signals was quantified using the AIDA software (Advanced Image Data Analyzer, Raytest) and normalized to β-actin intensity or Coomassie Blue staining as indicated.

**Tab. 9.4 List of Antibodies (AB) used for immunoblotting**

<b>1° AB</b>	<b>Antigen/Catalog No.</b>	<b>dilution</b>	<b>Source</b>
guinea pig-anti-Ae1	DCFIYEDQIRPQDREELLRALLK	1:500	Own (Pineda)
rabbit-anti-pendrin	CKDPLDLMEAEMNAEELDQDEAMRRLAS	1:1,000	Own (Pineda)
rabbit-anti-B1	CAQQDPASDTAL	1:1,000	Own (Pineda)
rabbit-anti-AQP2	AQP2 C-term	1:3,000	Loffing, J.
goat-anti-Foxi1	TSGVLYPREGTEV	1:250	abcam
goat-anti-Gdf15	sc46248	1:1,500	Santa Cruz
mouse-anti- $\beta$ actin	A5441	1:10,000	Sigma

<b>2° AB</b>	<b>Antigen/Catalog No.</b>	<b>dilution</b>	<b>Source</b>
anti-rabbit IgG, AP	S3731	1:5,000	Promega
anti-rabbit IgG, HRP	NA9341	1:10,000	Amersham
anti-guinea pig IgG, AP	A5062	1:5,000	Sigma
anti-mouse IgG, AP	S3721	1:5,000	Promega
anti-mouse IgG, HRP	NA931	1:10,000	Amersham
donkey-anti-goat IgG, AP	V1151	1:5,000	Promega

### **9.10 Blood and Urine Analysis**

Blood pH, blood gas ( $p\text{CO}_2$ ,  $p\text{O}_2$ ), and blood electrolyte concentrations ( $\text{Na}^+$ ,  $\text{K}^+$ ,  $\text{Cl}^-$ ,  $\text{Ca}^{++}$ ) were analyzed using an ABL 505 blood gas analyser (Radiometer, Copenhagen) and performed on heparinised venous blood as described earlier (chapter 9.1). Creatinine in urine was measured according to Jaffe's method [221]. Urinary ammonium was measured following Berthelot's protocol [222]. Plasma and urinary Urea levels were measured using diacetyl monoxime [223] and inorganic phosphate was determined using acid ammonium molybdate [224]. The urinary electrolytes were normalized to urinary creatinine and represented as mean  $\pm$  SEM (standard error of the mean).

### **9.11 Cell culture and transient transfection**

Hek293T cells (P8-10) were grown on 75 cm<sup>2</sup> flask (Greiner) in complete growth medium (DMEM with L-Glutamine (PAA), supplemented with 1 mM L-Glutamine (Lonza), 1 mM NEAA (Non-Essential-Amino-Acids (Lonza), and 10% FCS (fetal calf serum, Gibco)) at 37°C with 5% CO<sub>2</sub> atmosphere. Cells were passaged when they reached 80% confluency and split in a 1:15 ratio for general maintenance. Prior to trypsinization with 0.1 mM Trypsin-EDTA (Sigma), the monolayer was washed twice with prewarmed PBS (without Ca<sup>++</sup>, Mg<sup>++</sup>, Gibco). Trypsinization was stopped by addition of complete growth medium and cell suspension was centrifuged at 1,000 rpm for 1 minute. The supernatant was removed and the cell pellet was gently resuspended in complete growth medium. Transient transfections were carried out using Lipofectamine 2000 (Invitrogen) following manufacturer recommendations. Briefly, 2<sup>6</sup> cells/well were seeded on Poly-L-Lysine (Sigma) coated (15 minutes at RT) 6 well plates (Greiner) in complete growth medium. After 24 hours the growth medium was replaced by serum-free reduced medium (Opti-Mem, Gibco) and cells were transfected with 1 µg Foxi1 plasmid DNA (ImaGenes IRAVp968C0115D, Acc.No. BC007475). In case of untransfected cells, the plasmid DNA was replaced by equivalent amounts of sterile distilled water. Cells were harvested 48 hours after transfection and total cell lysates were prepared as described in chapter 9.7. Plasmid DNA was purified from a liquid bacteria culture using the Fast Plasmid Kit (Eppendorf).

### **9.12 Statistical Analysis**

All data are presented as mean ± SEM (standard error of the mean). Statistical Analysis was performed with GraphPad Prism Software 5.0 and the unpaired *t*-test or one way ANOVA with Tukey post-test were done where appropriate. Only results with  $p \leq 0.05$  were considered statistically significant.

## 10. RESULTS

As already mentioned, a urinary concentrating defect correlating with Aqp2 dysregulation was observed in adult Ae1 deficient mice, in addition to erythroidal defects and severe dRTA [190]. Furthermore, preliminary unpublished observations indicated the abnormal expression of renal IC marker proteins involved in acid-base homeostasis. Therefore, colocalization studies using specific antibodies targeting the  $\alpha 4$  and B1 subunits of the V-ATPase, Pds, Ae1, CaII and Aqp2 were performed. As described earlier, carbonic anhydrase II is strongly expressed in renal ICs and therefore commonly used as a general marker [41, 42]. However, in the present study a strong CaII expression was additionally found in Aqp2 positive cells of the CNT and early CCD in kidneys of murine and human origin (data not shown). Hence, CaII did not prove a specific marker for renal ICs in general and these data will not be further considered. Therefore, as shown in Fig. 2.6 renal principal cells were identified by the presence of Aqp2 (Aqp2<sup>+</sup>), while V-ATPase (B1<sup>+</sup>,  $\alpha 4$ <sup>+</sup>, V-ATPase<sup>+</sup>) labelling identified types of intercalated cells in general. Furthermore, Ae1 (Ae1<sup>+</sup>) expression specified type A-ICs, where appropriate, while Pds (Pds<sup>+</sup>) labelling determined type B-ICs. The number of cells positive or negative for either marker was assessed and statistically analyzed with respect to the segmental distribution. Within the CD system four different segments (CNT, CCD, OMCD and IMCD) were primarily distinguished based on morphologic criteria. Moreover, simultaneous immunologic localization of the Calcium binding protein Calbindin D28K (Calb1) helped to discriminate between the CNT and CCD segments. No further information was gathered by Calb1 staining and therefore these data will not be presented.

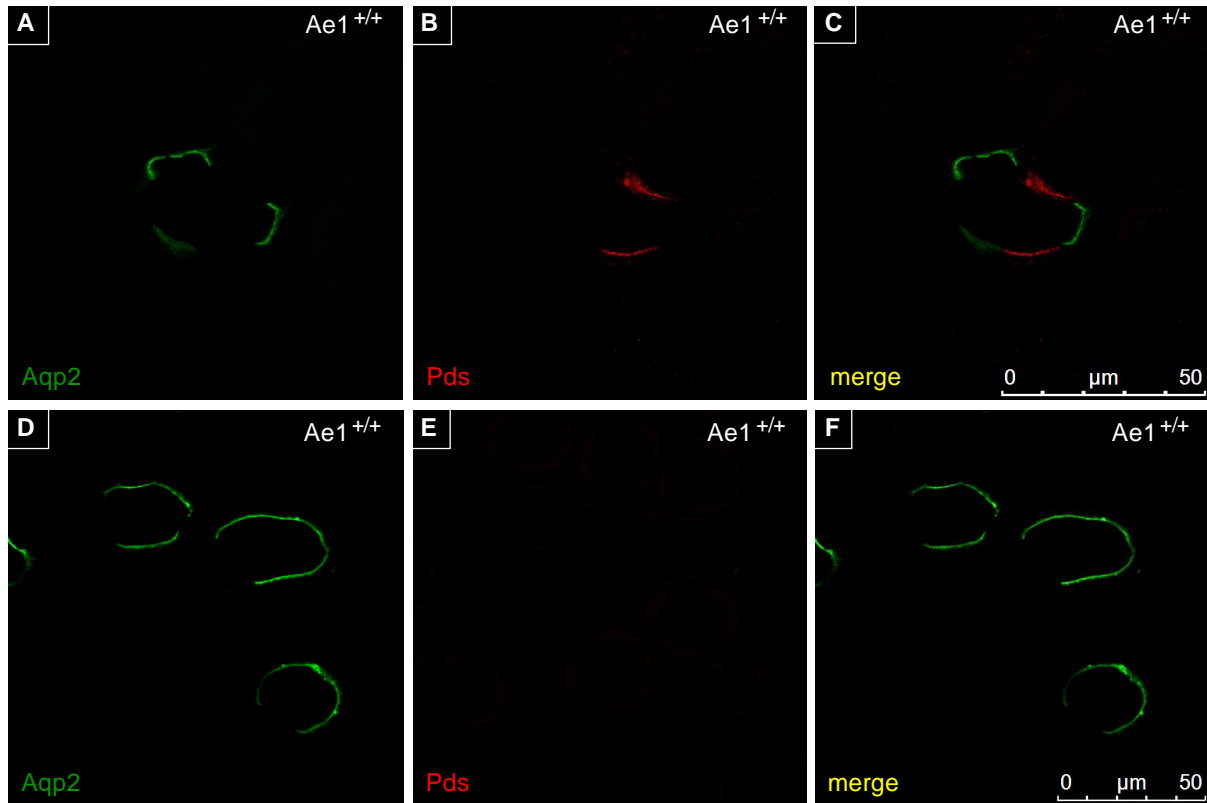
## ***10.1 Intercalated cells of the mature Ae1 deficient mouse kidney are abnormally differentiated***

Investigation of the mature kidney of juvenile (P30) and adult (P84) Ae1 deficient mice revealed unusual expression of pendrin in inner medullary collecting ducts. Moreover, mRNA expression levels as well as protein abundance and expression pattern of candidate regulatory molecules involved in terminal differentiation of the renal collecting duct were found to be abnormally regulated in kidneys of P30 and P84 Ae1 KO mice. However, kidney development and maturation seemed unaffected in Ae1 deficient animals.

### ***10.1.1 Pendrin is abnormally expressed in the inner medulla of adult Ae1 deficient mouse kidney***

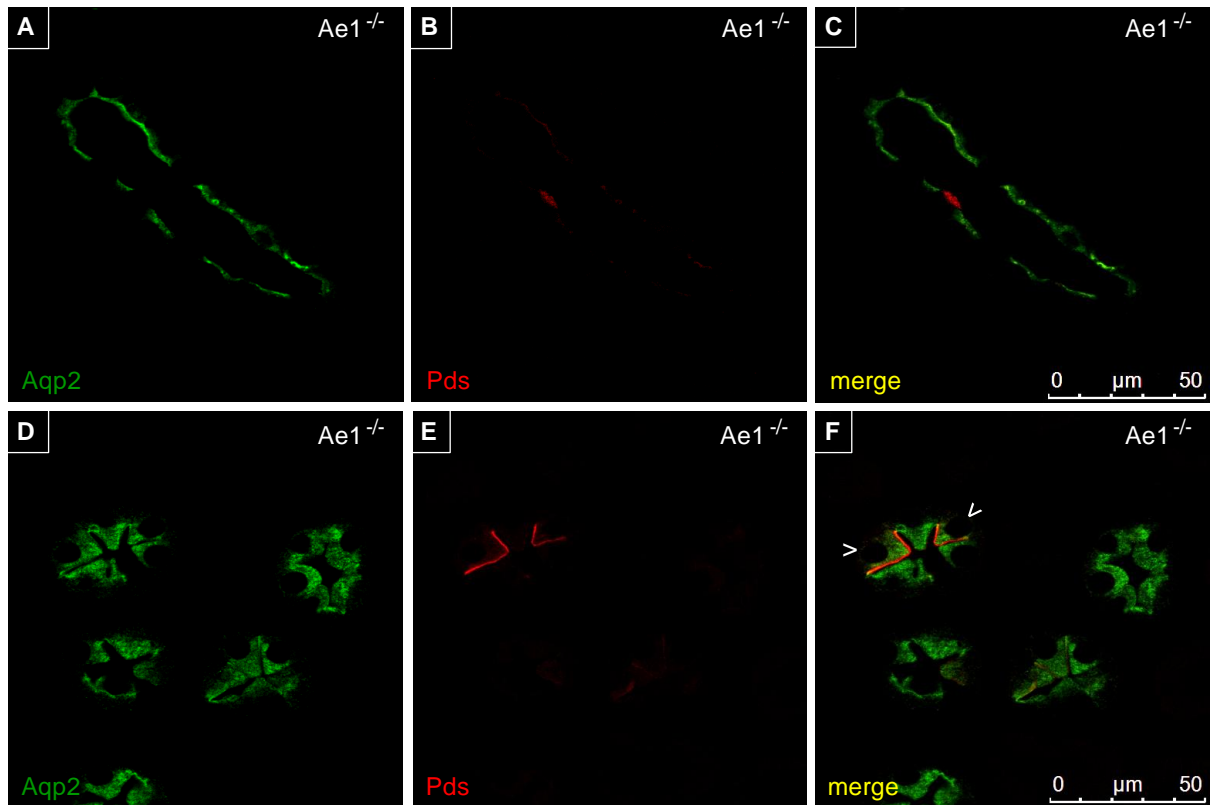
In the healthy adult kidney, pendrin expression identifies type B-ICs and is usually restricted to the renal cortex, (Fig. 10.1). In contrast to wild-type (WT), Pds expression was still present in the renal inner medulla of adult (P84, postnatal day 84) Ae1 knock-out (KO) mice (Fig. 10.2). Moreover, Pds in the inner medulla was unusually coexpressed with the PC marker Aqp2. Interestingly, Pds expression in the renal cortex (CNT and CCD) of adult Ae1 KO mice appeared normal, indicating a cortical preservation of the type B-IC phenotype. Nevertheless, quantification of individually labelled cells revealed an altered cellular distribution in distinct renal segments of Ae1 KO mice compared to WT (Fig. 10.3). The marker combination used in this colocalization study distinguishes only between principal cells (Aqp2<sup>+</sup>) and type B-ICs (Pds<sup>+</sup>), thus other variations of ICs are represented in the population of unlabelled (none) cells. Consistent with the chronic dRTA, the relative number of Pds<sup>+</sup> cells is significantly reduced in the CNT and CCD of Ae1 KO mice with a concomitant increase in Aqp2<sup>+</sup> cells (Fig. 10.3A, B). Furthermore, unusually Aqp2<sup>+</sup>/Pds<sup>+</sup> colabelled cells were detected in the renal OMCD (Fig. 10.3C), probably at the expense of unlabelled (none) cells. Moreover, the relative abundance of both, Pds<sup>+</sup> and

Aqp2<sup>+</sup>/Pds<sup>+</sup> cells respectively, is significantly increased in IMCDs of Ae1 deficient mice, with a simultaneous reduction of unlabeled cells (Fig. 10.3D).

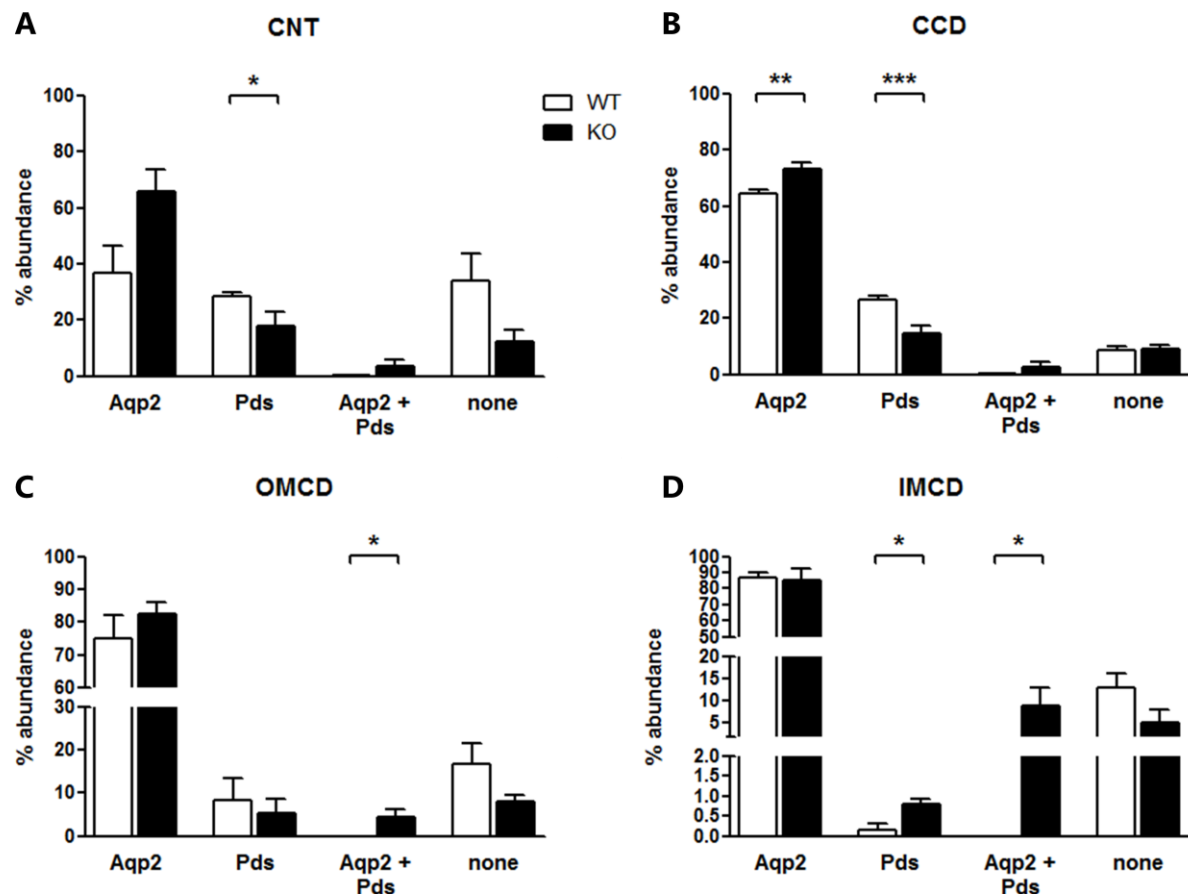


**Fig. 10.1 Immunologic localization of Pds in adult (P84) Ae1<sup>+/+</sup> mouse kidney. A-C)** Representative photomicrographs from renal cortex (CCD). **D-F)** Representative photomicrographs from renal inner medulla (IMCD). **A, D)** Individual Aqp2 (aquaporin 2) expression in renal CCD and IMCD identifies principal cells. **B)** In the CCD, Pds (pendrin) localizes to the luminal membrane of Aqp2 negative cells, identifying B-ICs. **E)** In adult WT mouse kidney, Pds is absent from the inner medulla. **C, F)** merge. Magnification 400x.





**Fig. 10.2 Immunologic localization of Aqp2 and Pds in adult (P84) *Ae1*<sup>-/-</sup> mouse kidney.** **A-C)** Representative photomicrographs from renal cortex (CCD). **D-F)** Representative photomicrographs from renal inner medulla (IMCD). **A, D)** Individual Aqp2 (aquaporin 2) expression in renal CCD and IMCD identifies principal cells. **B)** In the CCD, Pds (pendrin) localizes to the luminal membrane of Aqp2 negative cells, identifying B-ICs. **C)** merge. **E)** In adult *Ae1* deficient mouse kidney, Pds expression is occasionally still present on the luminal membrane of IMCD cells and even coexpressed with luminal Aqp2 expression (**F**, arrowheads). Magnification 400x.

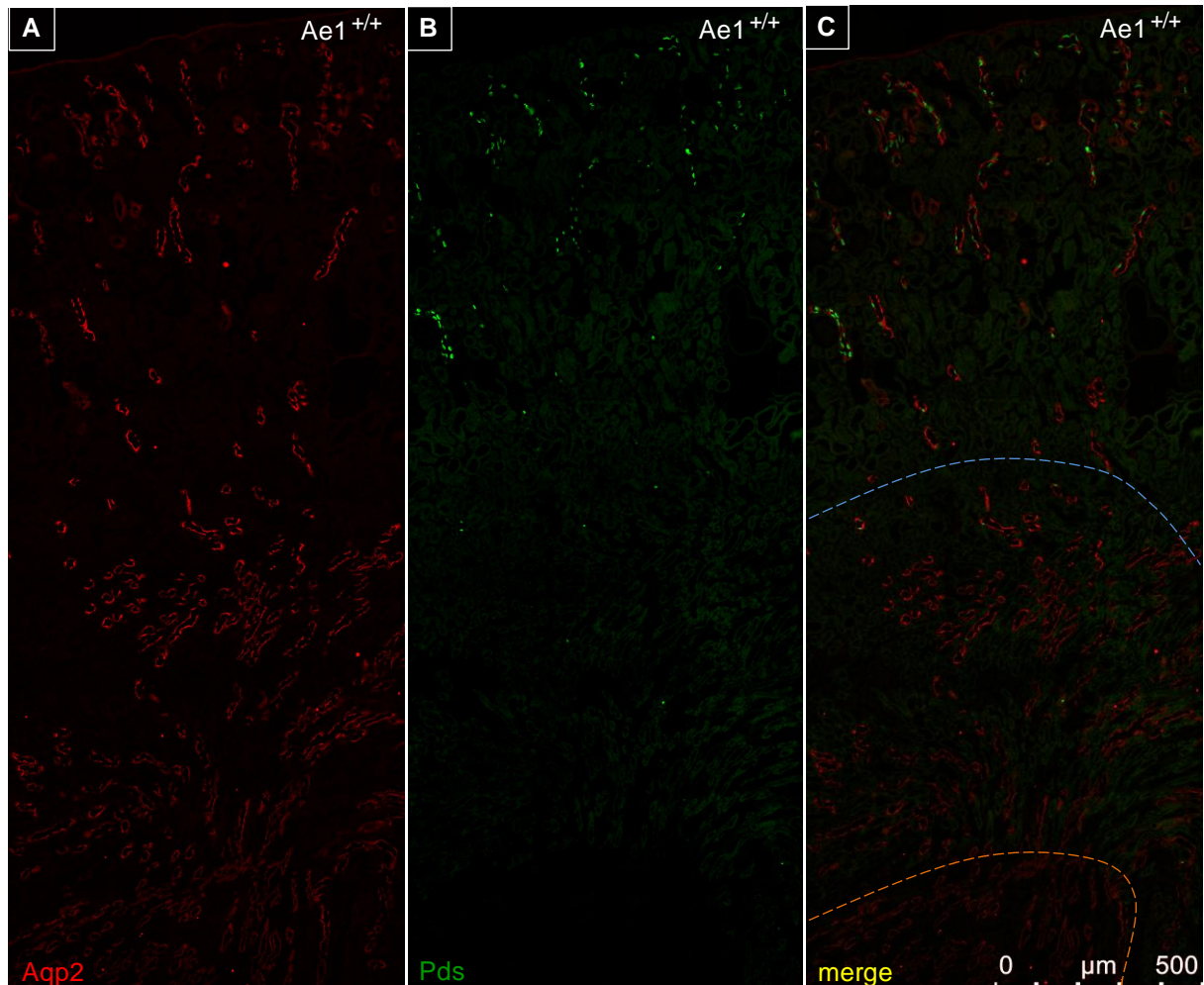


**Fig. 10.3 Relative distribution of Aqp2 and Pds in adult (P84) WT and Ae1 KO mouse kidney. A-D)** Summary of cell counts assessing the relative abundance of Aqp2 (aquaporin 2) and Pds (pendrin) positive cells in the renal CNT, CCD, OMCD and IMCD of adult (P84) Ae1 WT and KO mice. In total, 9,676 cells were counted in WT kidneys (n=7) and 5,533 cells in KO kidneys (n=4). Statistical analysis using Student's *t*-test. \*  $p \leq 0.05$ , \*\*  $p \leq 0.01$ , \*\*\*  $p \leq 0.001$ .  
CNT: connecting tubule, CCD: cortical collecting duct, OMCD: outer medullary collecting duct, IMCD: inner medullary collecting duct

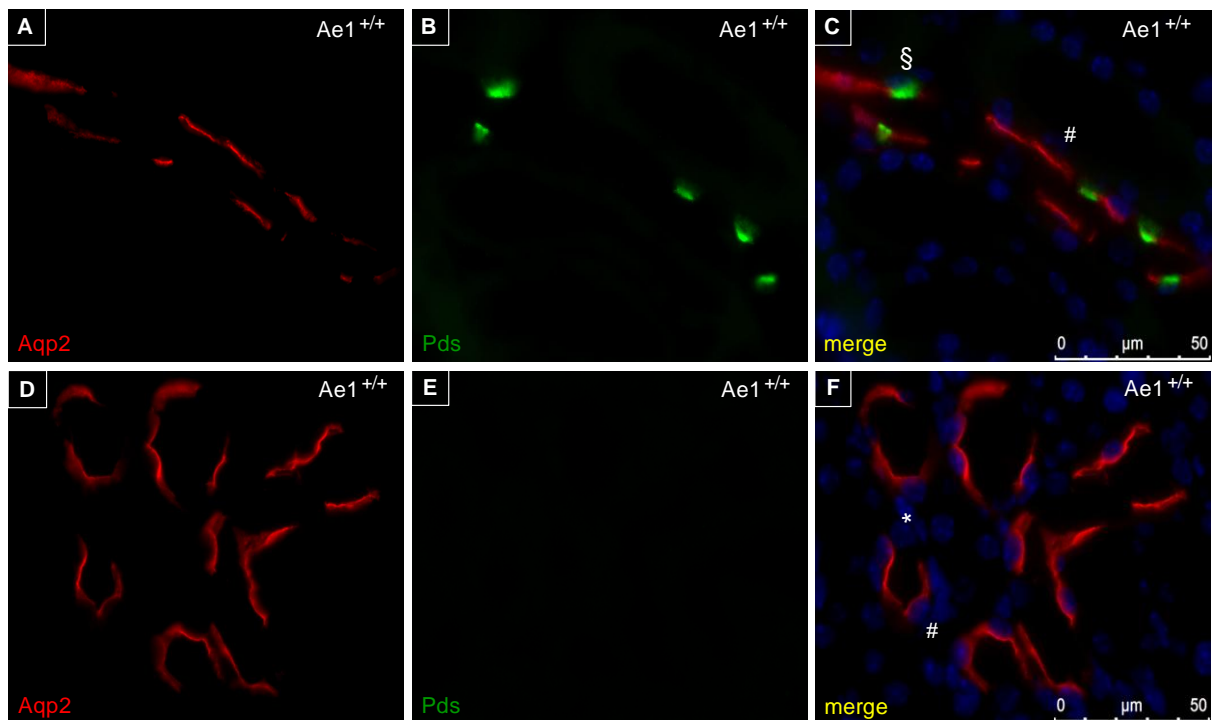
### ***10.1.2 Intercalated cells in the mature kidney of juvenile (P30) Ae1 deficient mice are abnormally differentiated***

Similar to adult (P84) Ae1 deficient mice, an unusual presence of Pds in cells of the mature inner medulla was observed in juvenile (P30) Ae1 KO mouse kidneys (Fig. 10.4 to 10.7). Consistently, renal medullary Pds expression was still present in Ae1 KO kidneys and in addition unusually coexpressed with Aqp2 in the IMCD (Fig. 10.6C, 10.7F, 10.8C, D). However, the relative distribution of labelled cells within the renal distal tubule was slightly different in P30 compared to P84 Ae1 KO kidney, possibly reflecting the impact of prolonged dRTA. In detail, Aqp2<sup>+</sup> cells were less abundant in the renal CNT of P30 WT and Ae1 KO mice compared to P84 CNT, while unlabelled cells were more abundant (Fig. 10.8A). Interestingly, the relative abundance of Pds<sup>+</sup> cells in the renal CNT was comparable at both ages. Furthermore and in contrast to P84 kidneys, there was no difference in the relative abundance of Aqp2<sup>+</sup>, Pds<sup>+</sup> or unlabelled cells in the CNT of P30 Ae1 WT and KO kidneys. Moreover, Pds<sup>+</sup> cells in the renal CCD of P30 Ae1 KO mice were increased compared to P30 WT, while the relative abundance of unlabelled cells was reduced and Aqp2<sup>+</sup> cells were not altered (Fig. 10.8B). In contrast, Pds<sup>+</sup> cells in the renal CCD of P84 Ae1 KO animals were significantly reduced, while unlabelled cells were not altered and Aqp2<sup>+</sup> cells were increased compared to P84 WT kidneys (Fig. 10.3B). Similar to the renal CCD of P30 animals, the relative abundance of Pds<sup>+</sup> cells in P30 Ae1 KO OMCD was increased when compared to P30 WT OMCD and the relative numbers of unlabelled cells were decreased (Fig. 10.8C). In contrast, OMCDs of P84 Ae1 KO mice displayed unusual presence of Aqp2<sup>+</sup>/Pds<sup>+</sup> cells compared to P84 WT, probably on the expense of unlabelled cells (Fig. 10.3C). Finally, in the renal IMCD of P30 Ae1 KO increased abundances of Pds<sup>+</sup> and Aqp2<sup>+</sup>/Pds<sup>+</sup> cells were found compared to P30 WT, with an adequate reduction of Aqp2<sup>+</sup> cells (Fig. 10.8D). Likewise, Pds<sup>+</sup> and Aqp2<sup>+</sup>/Pds<sup>+</sup> cells were increased in Ae1 KO IMCD of P84 mice compared to WT IMCD, but here the relative number of Aqp2<sup>+</sup> was not altered (Fig. 10.3D). The inappropriate increases in

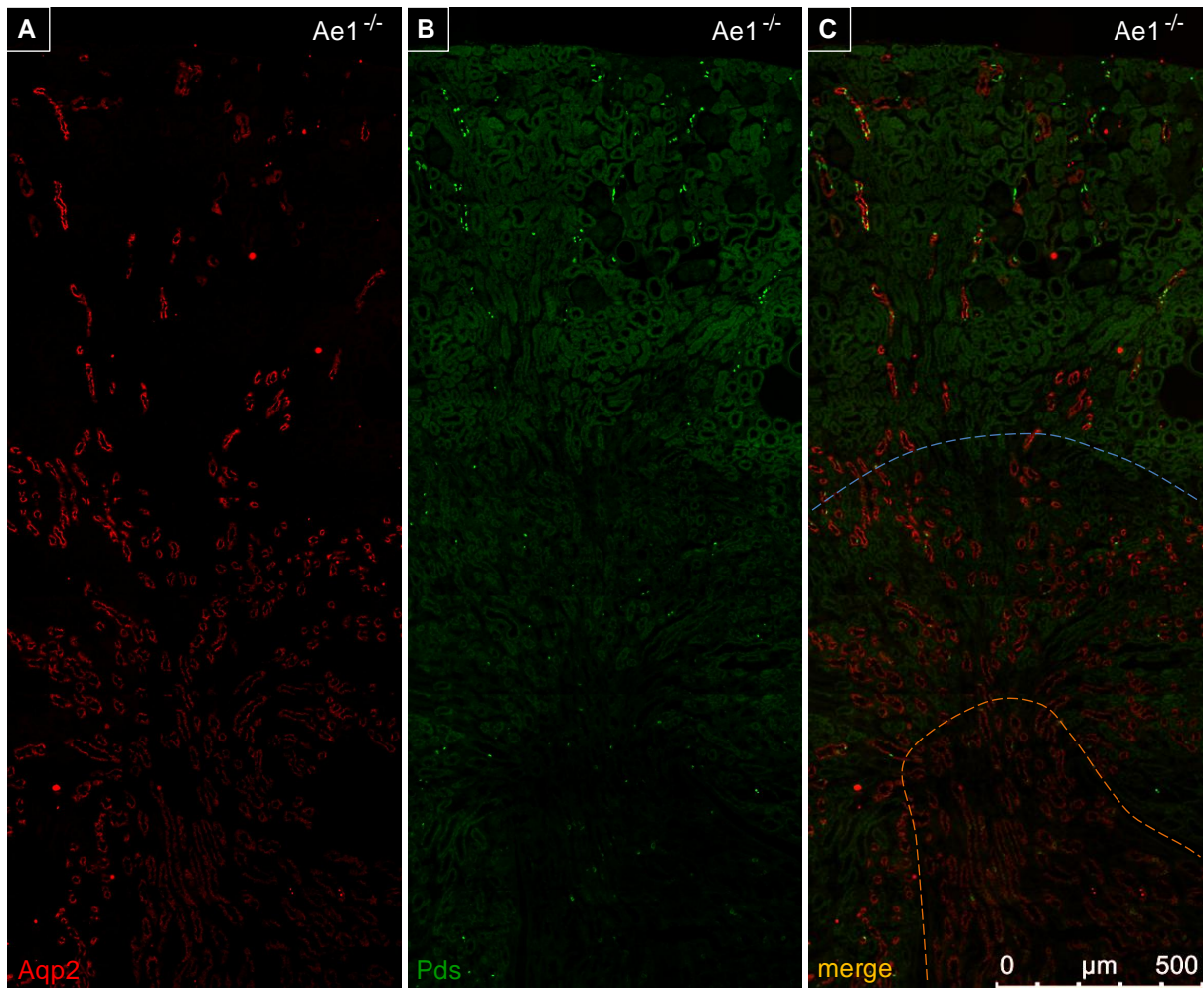
$Pds^+$  and  $Aqp2^+/Pds^+$  cells might rather be explained by the modestly reduced presence of unlabelled cells in P84 Ae1 KO kidneys compared to WT.



**Fig. 10.4 Zonal distribution of Aqp2 and Pds expressing cells in kidney of juvenile (P30) *Ae1*<sup>+/+</sup> mouse.** **A)** Aqp 2 (aquaporin 2) is expressed along cortical and medullary collecting ducts. **B)** Pds (pendrin) expression is restricted to the renal cortex in mature kidney. **C)** merge, blue dashed line indicating cortico-medullary boundary, orange dashed line indicating medullary-papillary boundary. Magnification 400x, Tile Scan.

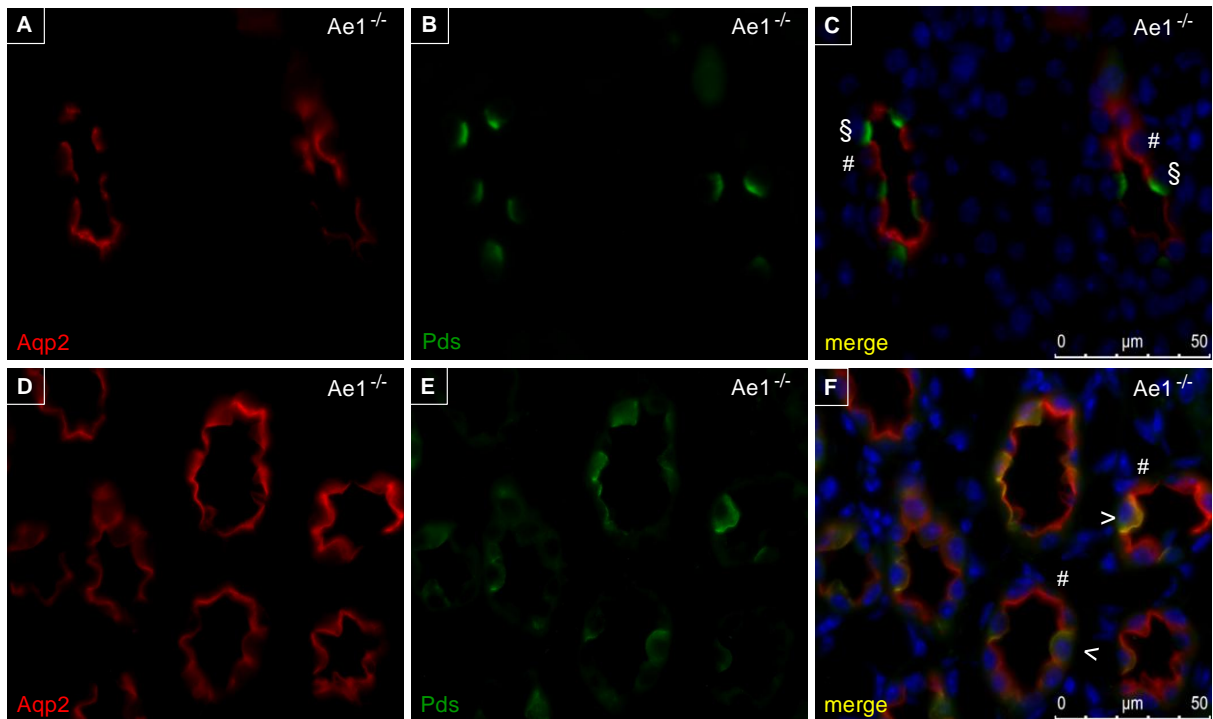


**Fig. 10.5 Zonal distribution of Aqp2 and Pds expressing cells in kidney of juvenile (P30) *Ae1*<sup>+/+</sup> mouse.** Aqp2 (aquaporin 2) is expressed along cortical (A) and medullary (D) collecting ducts. Pds (pendrin) expressing cells are frequently present in the renal cortex (B) and absent from the medullary collecting ducts (E). C, F merge. Nucleic acid stained with DAPI (blue). § Pds positive cell; # Aqp2 positive cell; \* Aqp2/Pds negative cell. Magnification 400x.

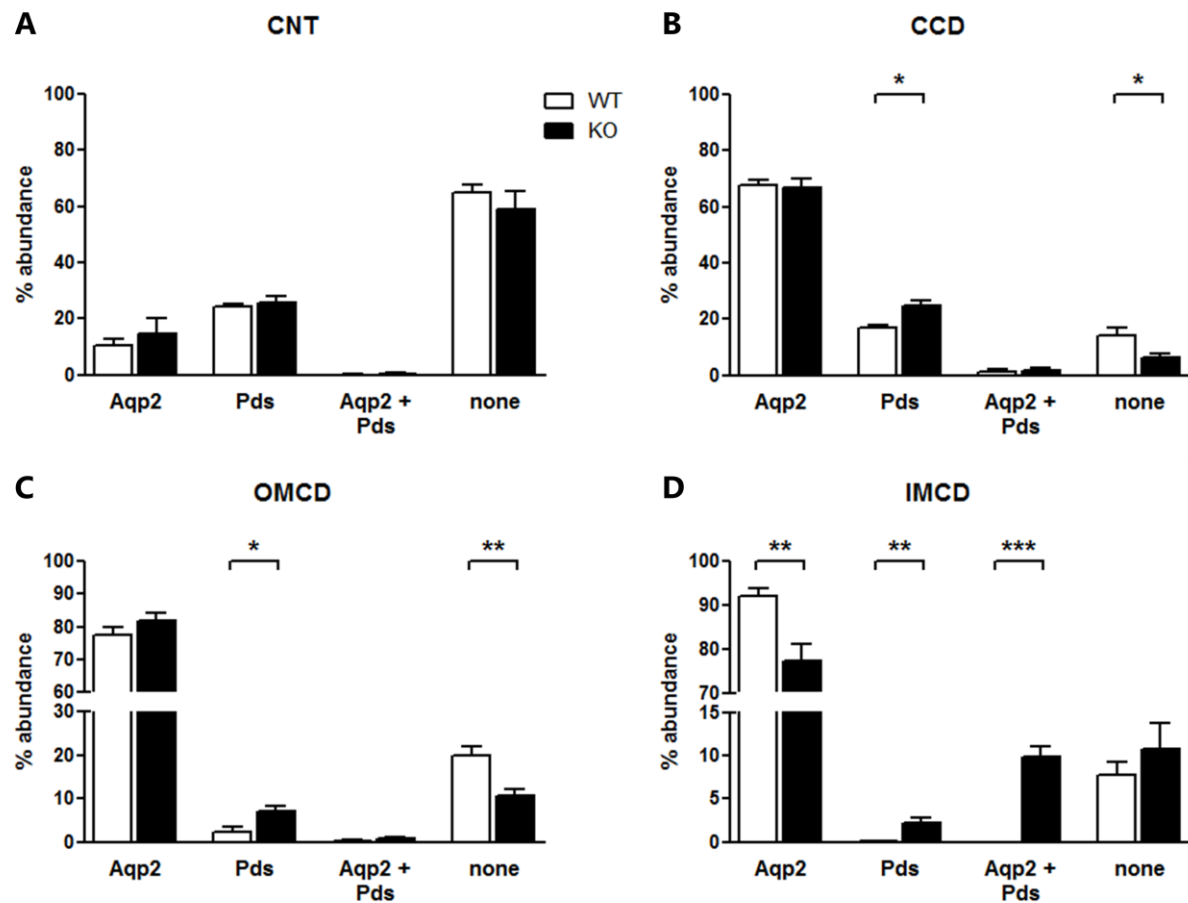


**Fig. 10.6 Zonal distribution of *Aqp2* and *Pds* expressing cells in kidney of juvenile (P30) *Ae1*<sup>-/-</sup> mouse.** **A)** Aqp 2 (aquaporin 2) is expressed along cortical and medullary collecting ducts. **B)** Pds (pendrin) expression is present in the cortex and inner medulla of the mature kidney. **C)** merge, blue dashed line indicating cortico-medullary boundary, orange dashed line indicating medullary-papillary boundary. Magnification 400x, Tile Scan.





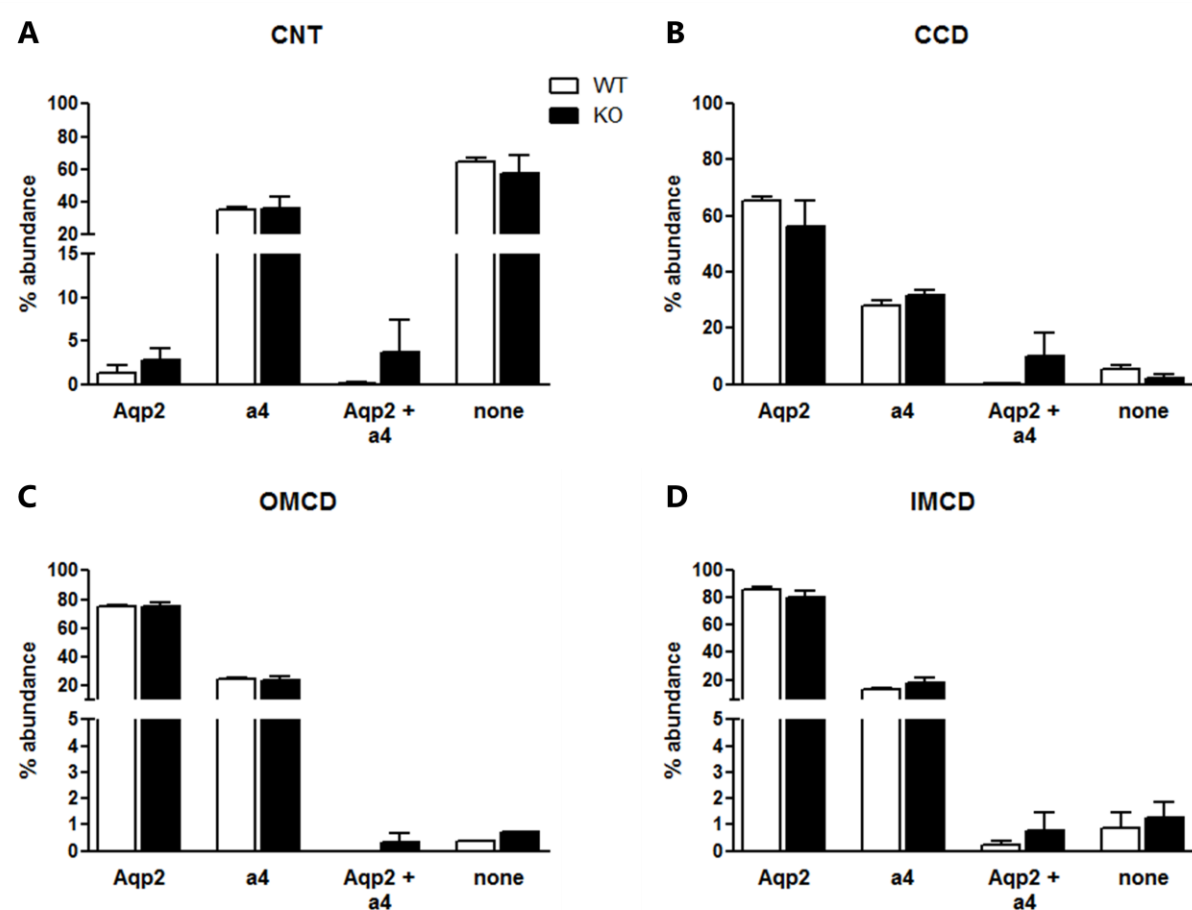
**Fig. 10.7 Zonal distribution of Aqp2 and Pds expressing cells in kidney of juvenile (P30) *Ae1*<sup>-/-</sup> mouse.** Aqp2 (aquaporin 2) is expressed along cortical (A) and medullary (D) collecting ducts. Pds (pendrin) expressing cells are frequently present in the cortical (B) and medullary collecting ducts (E). C, F merge. Nucleic acid stained with DAPI (blue). § Pds positive cell; # Aqp2 positive cell; arrowhead: Aqp2/Pds positive cell. Magnification 400x.



**Fig. 10.8 Relative distribution of Aqp2 and Pds expressing cells in kidneys of juvenile (P30) Ae1 WT and KO mice.** Summary of cell counts assessing the relative abundance of Aqp2 (aquaporin 2) and Pds (pendrin) positive cells in the renal CNT, CCD, OMCD and IMCD of juvenile (P30) Ae1 WT and KO mice. In total, 5,214 cells were counted in WT kidneys (n=5) and 5,335 cells in KO kidneys (n=5). Statistical analysis using Student's *t*-test. \*  $p \leq 0.05$ , \*\*  $p \leq 0.01$ , \*\*\*  $p \leq 0.001$ . CNT: connecting tubule, CCD: cortical collecting duct, OMCD: outer medullary collecting duct, IMCD: inner medullary collecting duct

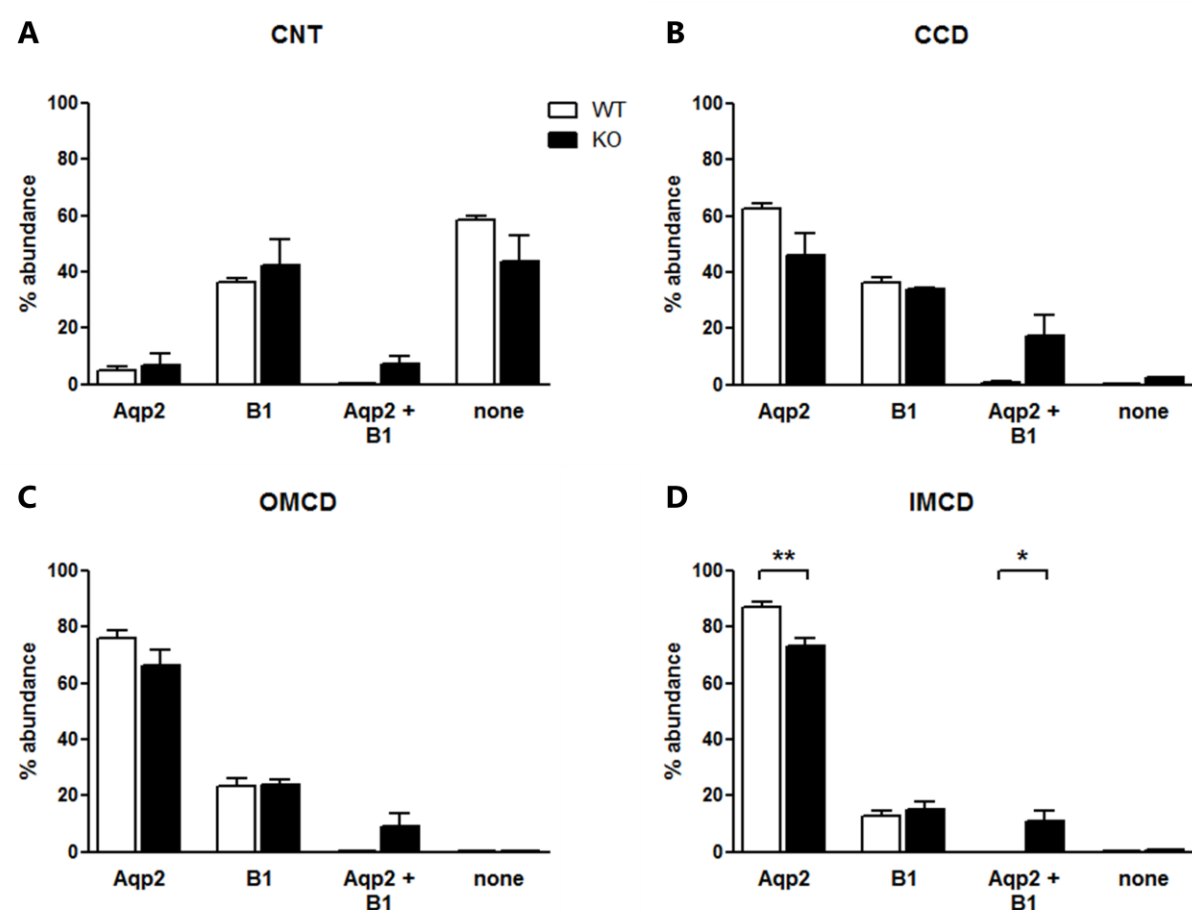


In order to investigate the distribution and appearance of Pds negative ICs of P30 Ae1 KO kidneys, co-immunostainings using the  $\alpha 4$  and B1 subunits of the V-ATPase, indicating the presence of manifestations of ICs other than type B-ICs, were performed and quantified (Fig. 10.9 and 10.10). No significant differences in the relative abundance and distribution of the kidney-specific  $\alpha 4$  subunit of V-ATPase ( $\alpha 4^+$ ) cells were found in P30 Ae1 KO kidneys compared to WT (Fig. 10.9). However, co-staining of Aqp2 and the IC-specific B1 subunit of V-ATPase gave another impression. Here, renal inner medullary collecting ducts of P30 Ae1 KO mice showed a significant increased abundance for  $\text{Aqp2}^+/\text{B1}^+$  cells with a concomitant reduction in  $\text{Aqp2}^+$  cells compared to WT (Fig. 10.10D). Although alterations of  $\text{B1}^+$  cell expression in the other segments did not reach significant difference, a similar trend was observed. In the renal CNT of P30 Ae1 KO mice a modest increase in  $\text{Aqp2}^+/\text{B1}^+$  cells (p-value 0.0590) and reduction in unlabelled cells became apparent, while the increase in  $\text{Aqp2}^+/\text{B1}^+$  cells of the CCD (p-value 0.0621) and OMCD (p-value 0.0798) was accompanied by a reduction in  $\text{Aqp2}^+$  cells (Fig. 10.10A-C).



**Fig. 10.9 Relative distribution of Aqp2 and a4 expressing cells in kidneys of juvenile (P30) Ae1 WT and KO mice.** Summary of cell counts assessing the relative abundance of Aqp2 (aquaporin 2) and a4 (a4 subunit of V-ATPase, Atp6v0a4) positive cells in the renal CNT, CCD, OMCD and IMCD of juvenile (P30) Ae1 WT and KO mice. In total, 4,930 cells were counted in WT kidneys (n=5) and 5,094 cells in KO kidneys (n=5). Statistical analysis using Student's *t*-test. \*  $p \leq 0.05$ , \*\*  $p \leq 0.01$ , \*\*\*  $p \leq 0.001$ .

CNT: connecting tubule, CCD: cortical collecting duct, OMCD: outer medullary collecting duct, IMCD: inner medullary collecting duct



**Fig. 10.10 Relative distribution of Aqp2 and B1 subunit of V-ATPase (*Atp6v1b1*) expressing cells in kidneys of juvenile (P30) *Ae1* WT and KO mice.** Summary of cell counts assessing the relative abundance of Aqp2 (aquaporin 2) and B1 (B1 subunit of V-ATPase, *Atp6v1b1*) positive cells in the renal CNT, CCD, OMCD and IMCD of juvenile (P30) *Ae1* WT and KO mice. In total, 5,619 cells were counted in WT kidneys (n=5) and 4,638 cells in KO kidneys (n=5). Statistical analysis using Student's *t*-test.

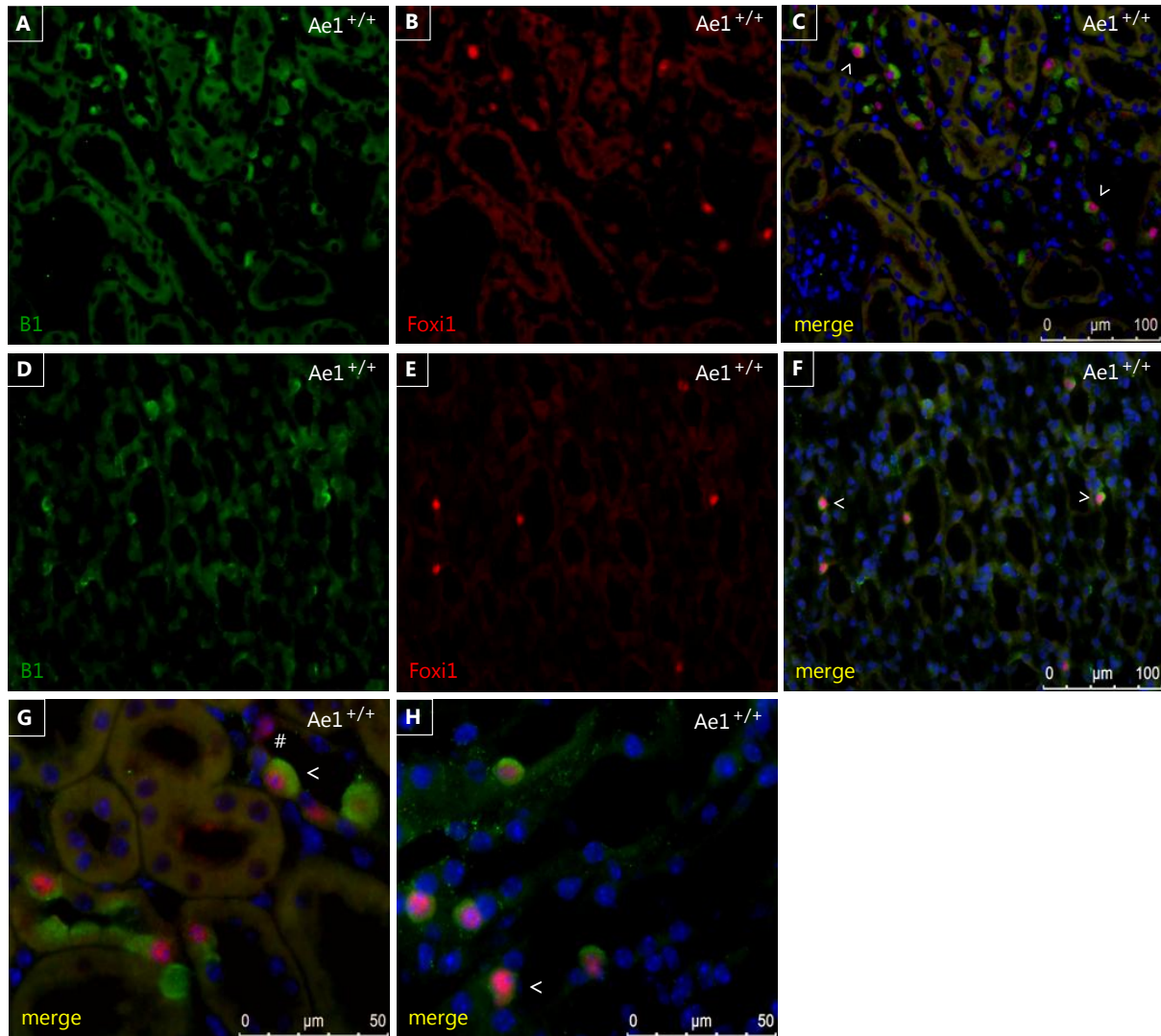
\*  $p \leq 0.05$ , \*\*  $p \leq 0.01$ , \*\*\*  $p \leq 0.001$ .

CNT: connecting tubule, CCD: cortical collecting duct, OMCD: outer medullary collecting duct, IMCD: inner medullary collecting duct

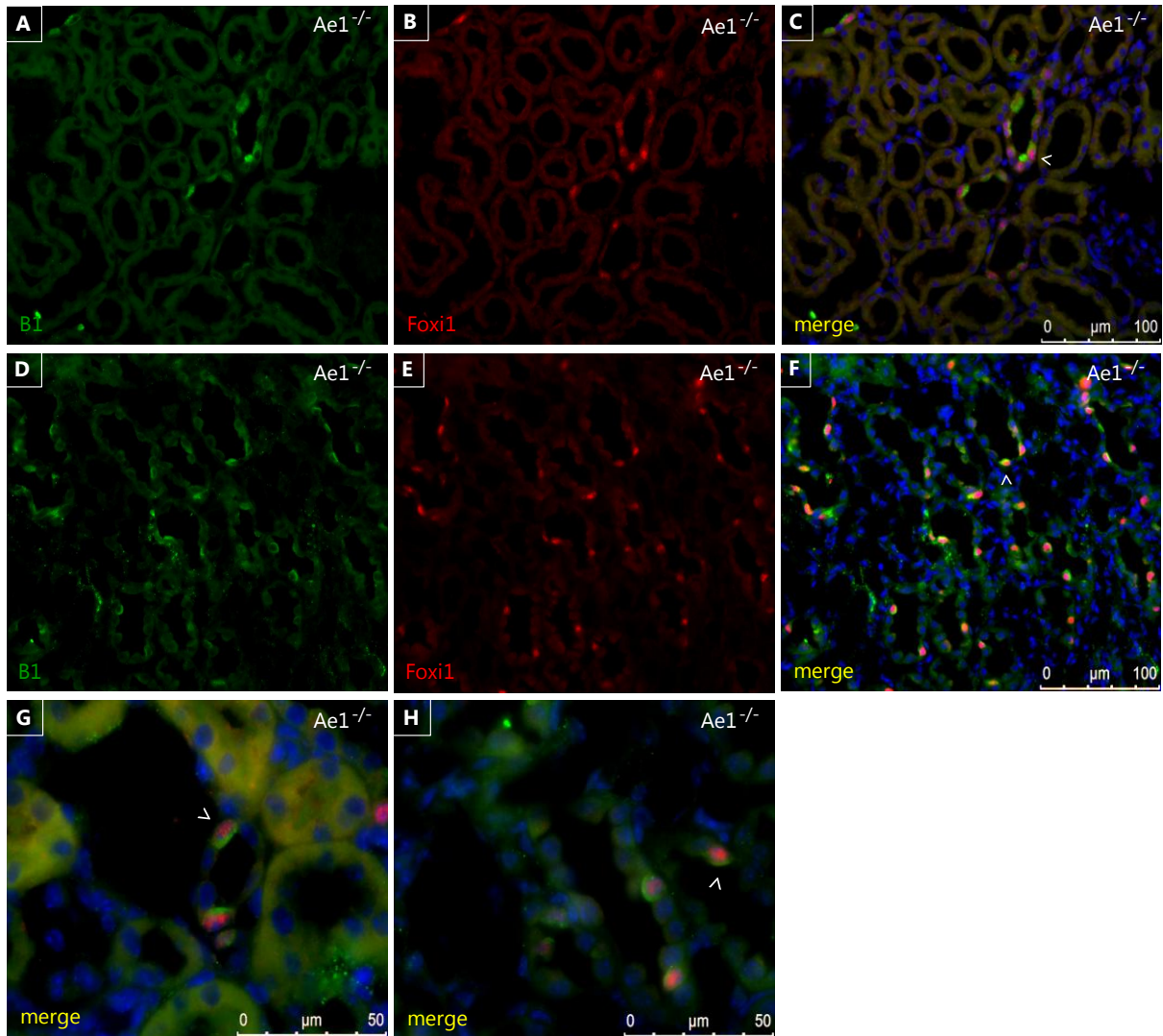
As described in chapter 2.7, some molecules and transcription factors have been identified as critical in terminal differentiation of renal CD specific cells. Among them are the transcription factors Foxi1 and Cp2l1. Even though both transcription factors were localized in renal tubular structures a detailed localization concerning their putative presence in the CD epithelium is missing. Therefore, the expression patterns of both transcription factors were investigated in more detail and preliminary results were compared between Ae1 WT and KO.

The Foxi1 protein was occasionally detected in nuclei of CD cells in P30 WT mouse kidney. Co-labelling with the B1 subunit of V-ATPase (Fig. 10.11, 10.13) and Aqp2 (Fig. 10.14, 10.16) revealed predominant expression of Foxi1 in intercalated cells and less in principal cells of the CNT, CCD, OMCD and IMCD. Quantification of cell counts showed that the majority of Foxi1<sup>+</sup> nuclei corresponded with B1<sup>+</sup> cells (Fig. 10.13, 10.16). Results obtained from cell counts suggest a relative equal distribution of Foxi1<sup>+</sup>/B1<sup>+</sup> cells among the CNT (23.69%), CCD (26.33%) and OMCD (21.70%) segments of P30 WT mouse kidney with a similar trend in Ae1 KO animals (Fig. 10.13). However, in Ae1 WT kidney the relative abundance of Foxi1<sup>+</sup>/B1<sup>+</sup> cells in the IMCD (15.19%) is significantly lower compared to CCD (26.33%, p-value <0.05), reflecting the expected reduced presence of ICs in the renal inner medulla compared to cortex. Similarly, in Ae1 WT kidney the presence of Foxi1<sup>+</sup> cells in the staining series with Aqp2 (Fig. 10.16) declines significantly from CNT (24.08%), CCD (18.64%), OMCD (14.72%) to IMCD (8.19%), indicating again the expected reduction of ICs along the CD system towards the papilla. Anyhow, comparing the relative distribution of Foxi1 expression in Ae1 WT and KO kidneys of P30 mice showed some small but significant differences. Individually labelled Foxi1<sup>+</sup> cells in the staining series with the B1 subunit of V-ATPase were significantly increased in IMCDs of Ae1 KO kidneys when compared to WT (Fig. 10.13D). In addition, a simultaneous but statistically not significant increase in Foxi1<sup>+</sup>/Aqp2<sup>+</sup> cells was observed in IMCDs of Ae1 KO kidneys (Fig. 10.16D). Furthermore, individually Foxi1<sup>+</sup> and unlabelled (none) cells were significantly reduced in the CCD of P30 Ae1 KO kidneys, while Aqp2<sup>+</sup>/Foxi1<sup>+</sup>

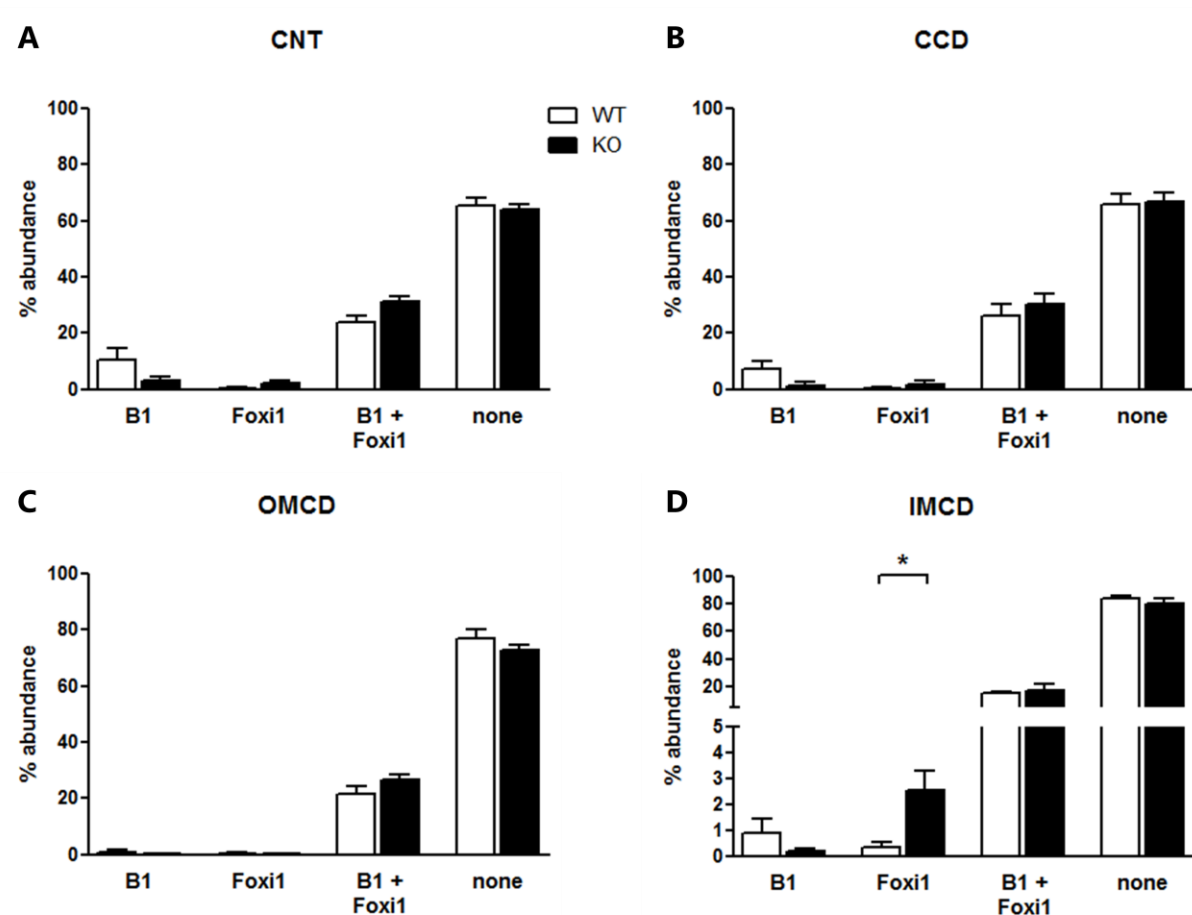
coexpressing cells increased concomitantly (Fig. 10.16B). Hence, loss of Ae1 in kidneys of P30 mice alters the relative segmental distribution of Foxi1 in the CCD and OMCD, favouring its coexpression with the principal cell marker Aqp2.



**Fig. 10.11 Expression pattern of Foxi1 and B1 expressing cells in kidney of juvenile (P30) *Ae1*<sup>+/+</sup> mouse.** **A-C, G)** Renal cortex. **D-F, H)** Renal inner medulla. B1 (B1 subunit of V-ATPase, Atp6v1b1) expressing intercalated cells were frequently present in the renal cortical (**A**) and medullary (**D**) collecting ducts. Nuclear expression pattern of Foxi1 protein was detected in cortical (**B**) and medullary (**E**) tubular structures. Overlay reveals expression of Foxi1 in B1 positive and less frequently in B1 negative cells along the renal collecting ducts (**C, F, G, H**). Nucleic acid stained with DAPI (blue). # B1 negative/Foxi1 positive cell; arrowhead: B1/Foxi1 positive cell. **A-F)** Magnification 400x. **G,H)** Magnification 1,000x.



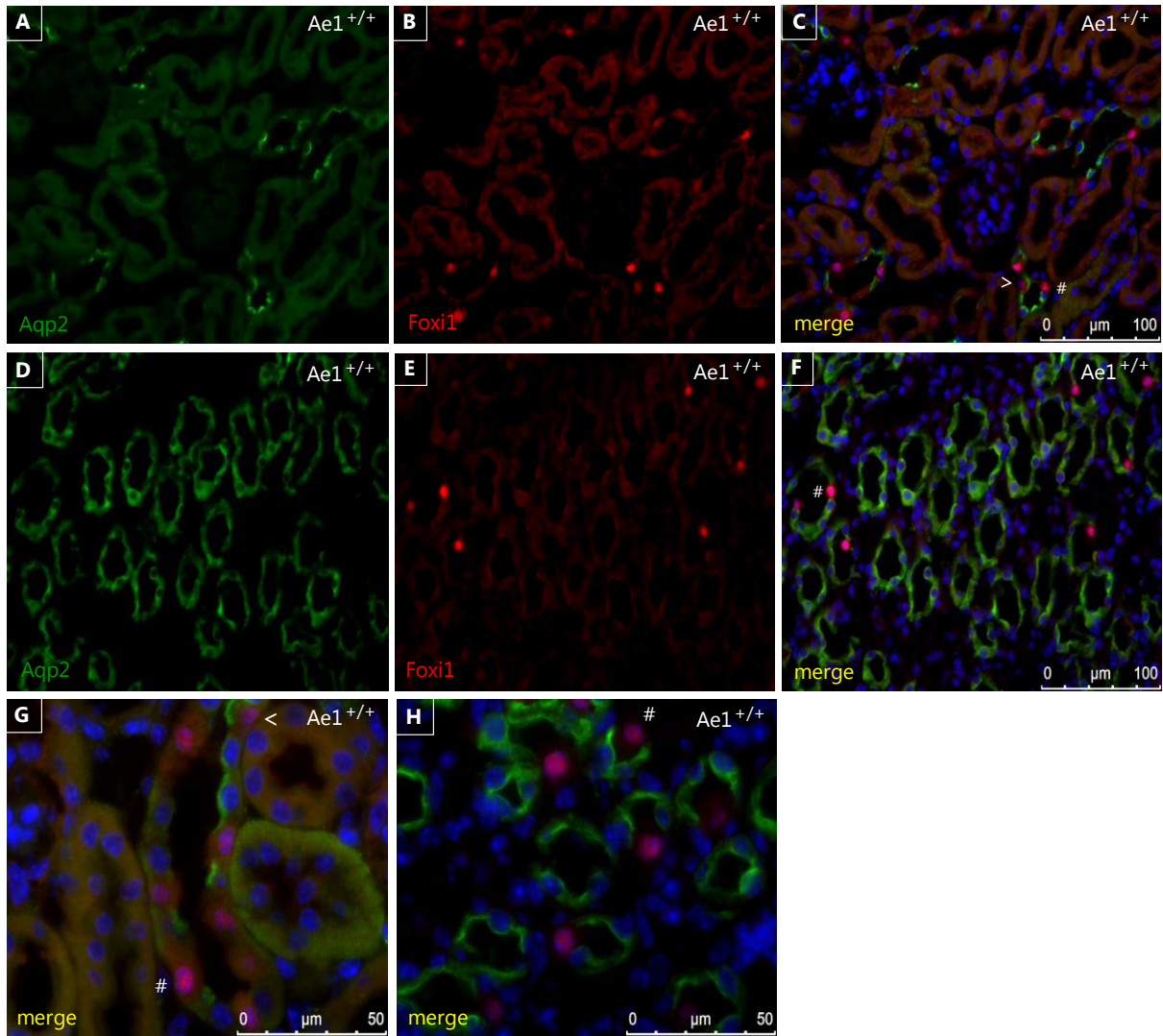
**Fig. 10.12 Expression pattern of Foxi1 and B1 expressing cells in kidney of juvenile (P30) *Ae1*<sup>-/-</sup> mouse.** **A-C, G)** Renal cortex. **D-F, H)** Renal inner medulla. B1 (B1 subunit of V-ATPase, Atp6v1b1) expressing intercalated cells were frequently present in the renal cortical (**A**) and medullary (**D**) collecting ducts. Nuclear expression pattern of Foxi1 protein was detected in cortical (**B**) and medullary (**E**) tubular structures. Overlay reveals expression of Foxi1 in B1 positive and less frequently in B1 negative cells along the renal collecting ducts (**C, F, G, H**). Nucleic acid stained with DAPI (blue). arrowhead: B1/Foxi1 positive cell. **A-F)** Magnification 400x. **G,H)** Magnification 1,000x.



**Fig. 10.13 Relative distribution of Foxi1 and B1 subunit of V-ATPase (*Atp6v1b1*) expressing cells in kidneys of juvenile (P30) *Ae1* WT and KO mice.** Summary of cell counts assessing the relative abundance of Foxi1 and B1 (B1 subunit of V-ATPase, *Atp6v1b1*) positive cells in the renal CNT, CCD, OMCD and IMCD of juvenile (P30) *Ae1* WT and KO mice. In total, 2,680 cells were counted in WT kidneys (n=4) and 3,489 cells in KO kidneys (n=5). Statistical analysis using Student's *t*-test. \*  $p \leq 0.05$ , \*\*  $p \leq 0.01$ , \*\*\*  $p \leq 0.001$ .

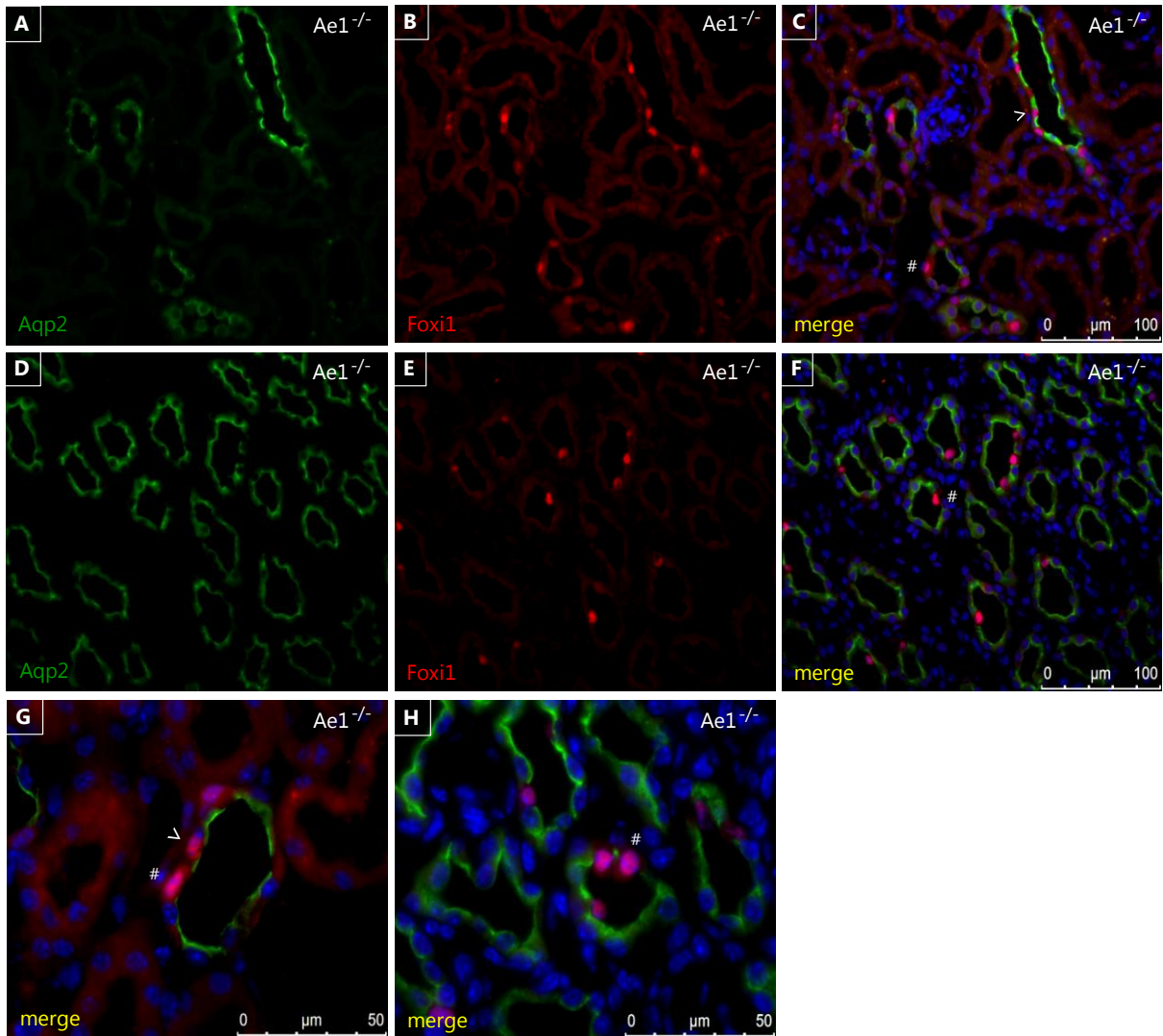
CNT: connecting tubule, CCD: cortical collecting duct, OMCD: outer medullary collecting duct, IMCD: inner medullary collecting duct



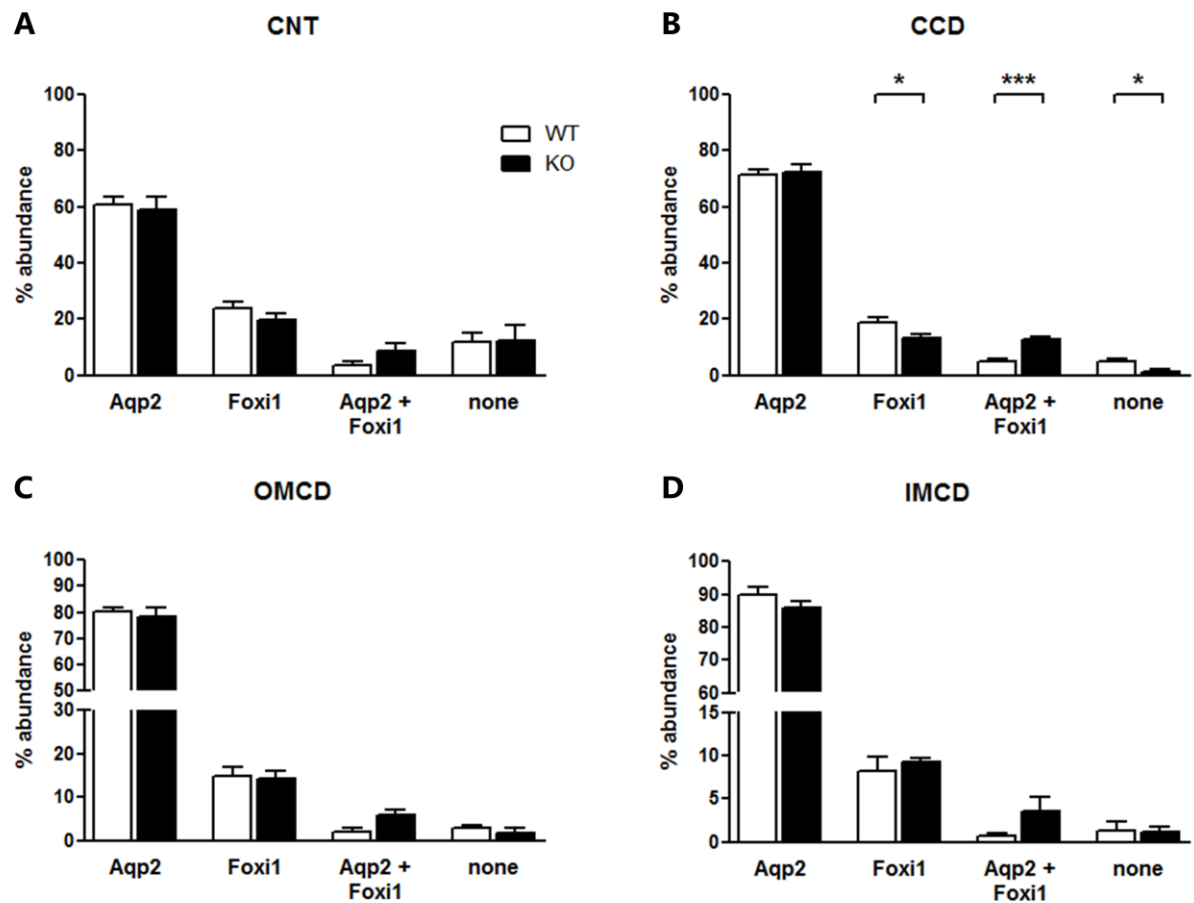


**Fig. 10.14 Expression pattern of Foxi1 and Aqp2 expressing cells in kidney of juvenile (P30) *Ae1*<sup>+/+</sup> mouse.** **A-C, G)** Renal cortex. **D-F, H)** Renal inner medulla. Principal cells expressing luminal Aqp2 (aquaporin 2) in renal cortical (**A**) and medullary (**D**) collecting ducts. Nuclear expression pattern of Foxi1 protein was detected in cortical (**B**) and medullary (**E**) tubular structures. Overlay reveals predominant expression of Foxi1 in Aqp2 negative and less frequently in Aqp2 positive cells along the renal collecting ducts (**C, F, G, H**). Nucleic acid stained with DAPI (blue). # Aqp2 negative/Foxi1 positive cell; arrowhead: Aqp2/Foxi1 positive cell. **A-F)** Magnification 400x. **G, H)** Magnification 1,000x.





**Fig. 10.15 Expression pattern of Foxi1 and Aqp2 expressing cells in kidney of juvenile (P30) *Ae1*<sup>-/-</sup> mouse.** **A-C, G)** Renal cortex. **D-F, H)** Renal inner medulla. Principal cells expressing luminal Aqp2 (aquaporin 2) in renal cortical (**A**) and medullary (**D**) collecting ducts. Nuclear expression pattern of Foxi1 protein was detected in cortical (**B**) and medullary (**E**) tubular structures. Overlay reveals predominant expression of Foxi1 in Aqp2 negative and less frequently in Aqp2 positive cells along the renal collecting ducts (**C, F, G, H**). Nucleic acid stained with DAPI (blue). #: Aqp2 negative/Foxi1 positive cell. arrowhead: Aqp2/Foxi1 positive cell. **A-F)** Magnification 400x. **G, H)** Magnification 1,000x.

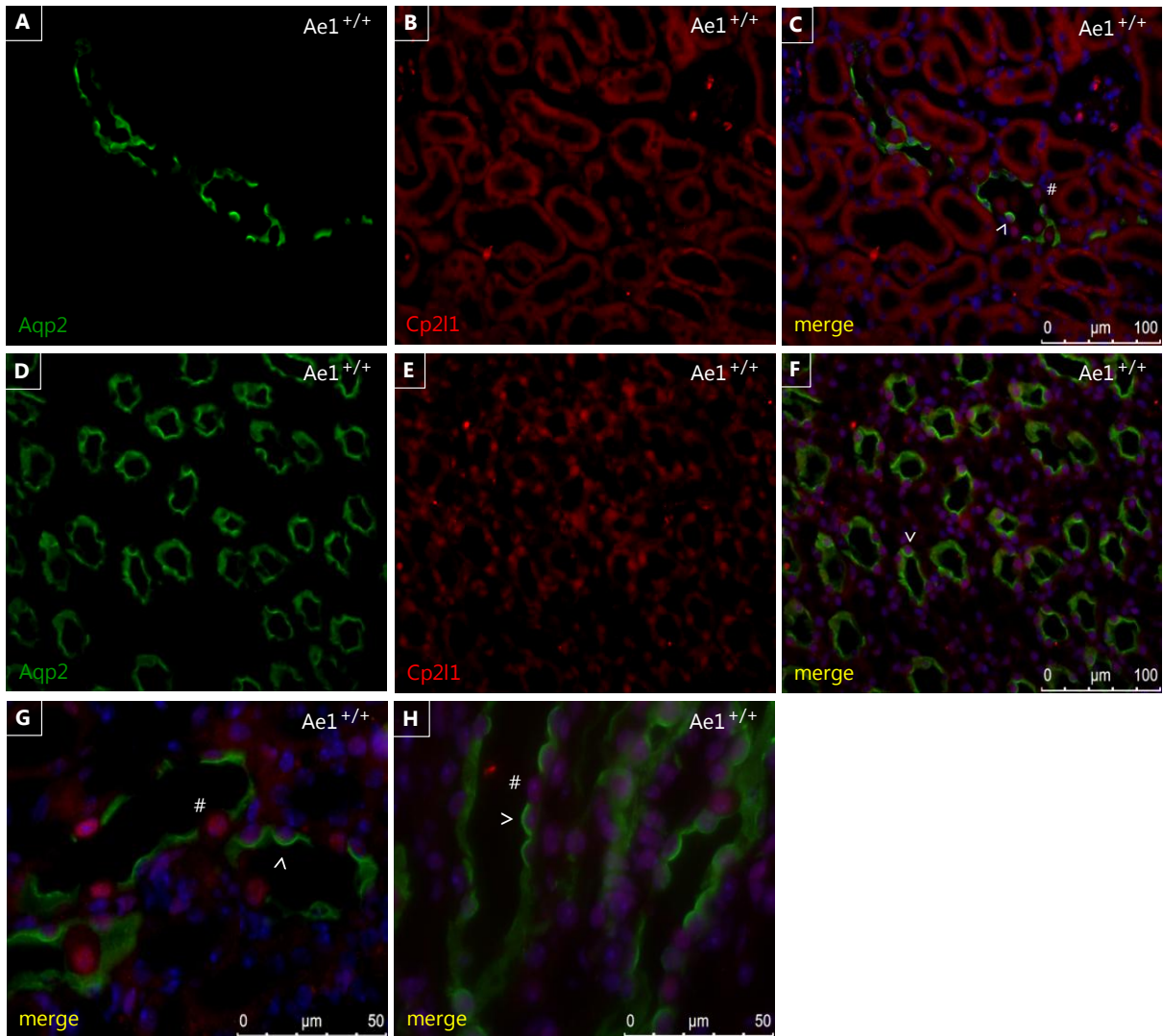


**Fig. 10.16 Relative distribution of Foxi1 and Aqp2 expressing cells in kidneys of juvenile (P30)**

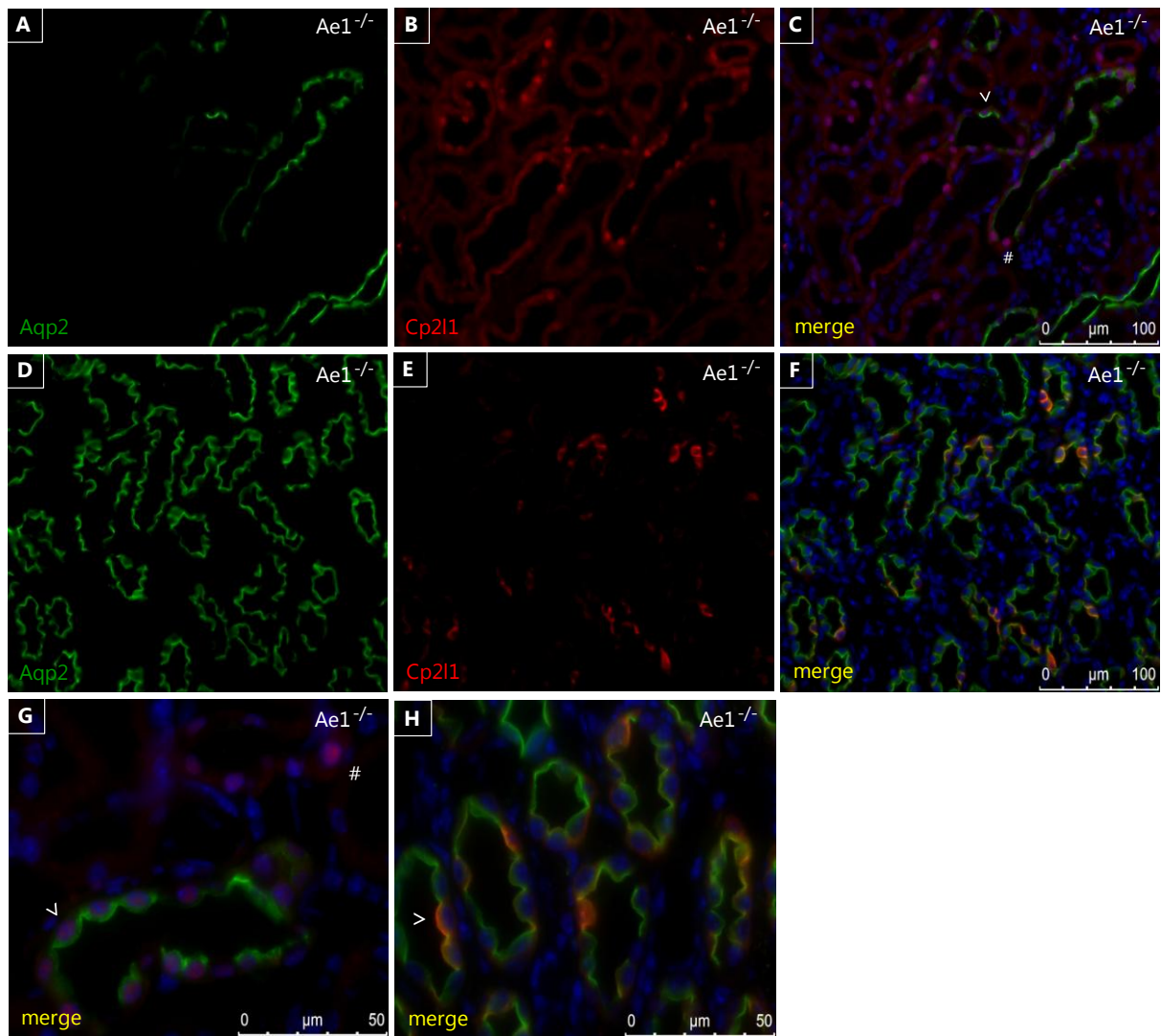
**Ae1 WT and KO mice.** Summary of cell counts assessing the relative abundance of Foxi1 and Aqp2 (aquaporin 2) positive cells in the renal CNT, CCD, OMCD and IMCD of juvenile (P30) Ae1 WT and KO mice. In total, 4,403 cells were counted in WT kidneys (n=5) and 5,243 cells in KO kidneys (n=5). Statistical analysis using Student's *t*-test. \*  $p \leq 0.05$ , \*\*  $p \leq 0.01$ , \*\*\*  $p \leq 0.001$ .

CNT: connecting tubule, CCD: cortical collecting duct, OMCD: outer medullary collecting duct, IMCD: inner medullary collecting duct

The expression of Cp2l1 was reported to change in accordance with the maturation of the developing nephron and CD system (Fig. 2.12) [90]. In adult mouse kidney, Cp2l1 expression was reported in parts of the DCT, CNT, CCD, and OMCD (Fig. 2.12). However, the cell types expressing Cp2l1 were not further characterized in this study [90]. In the present study, Cp2l1 expression was assessed in P30 kidneys and consistent with previous reports occasionally detected in cells of the DCT, CNT, CCD, and OMCD. Additionally, Cp2l1 was present in cells of the TAL and IMCD. In order to determine the cell type harbouring nucleic Cp2l1, coimmunostainings using the principal cell marker Aqp2 were performed. Interestingly, Cp2l1 was occasionally expressed in Aqp2 positive tubules of the renal cortex and medulla of P30 Ae1 WT (Fig. 10.17) and KO (Fig. 10.18). Moreover, Cp2l1 expression was detected in principal cells (Aqp2<sup>+</sup>) and manifestations of intercalated cells (Aqp2<sup>-</sup>) along the distal tubule. Strikingly, Cp2l1 was found to be present in virtually all nuclei of the IMCD in P30 WT kidneys (Fig. 10.17E, F, H), whereas the nuclear localization was completely absent in IMCDs of P30 Ae1 KO kidney (Fig. 10.18E, F, H). Moreover, instead of the expected nuclear pattern, the signal for Cp2l1 was now detected on the luminal cell membrane of certain cells, but not in all. However, detailed quantification using distinct CD markers are necessary to further analyze the expression pattern of Cp2l1. Furthermore, while analyzing the immunofluorescent images of Cp2l1, the relative signal intensity for Cp2l1, particularly in the renal cortex, appeared stronger in Ae1 deficient kidneys compared to WT (Fig. 10.17B, 10.18B). However, at present immunoblotting of Cp2l1 to assess the relative protein abundance was not successful due to technical reasons.



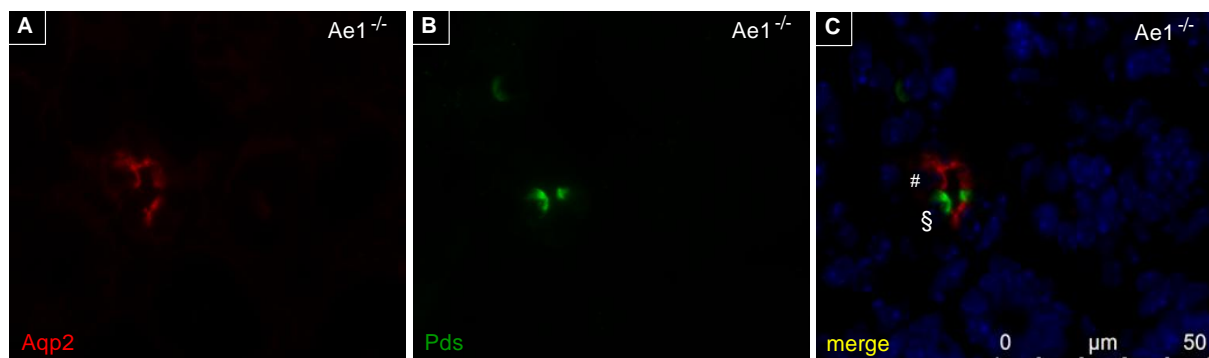
**Fig. 10.17 Expression pattern of *Cp2l1* and *Aqp2* expressing cells in kidney of juvenile (P30) *Ae1*<sup>+/+</sup> mouse.** **A-C, G)** Renal cortex. **D-F, H)** Renal inner medulla. Principal cells expressing luminal Aqp2 (aquaporin 2) in renal cortical (**A**) and medullary (**D**) collecting ducts. Nuclear expression pattern of Cp2l1 protein was detected in cortical (**B**) and predominantly medullary (**E**) tubular structures. Overlay reveals expression of Cp2l1 in Aqp2 positive and negative cells along the renal collecting ducts (**C, F, G, H**). Nucleic acid stained with DAPI (blue). # Aqp2 negative/Cp2l1 positive cell; arrowhead: Aqp2/Cp2l1 positive cell. **A-F)** Magnification 400x. **G, H)** Magnification 1,000x.



**Fig. 10.18 Expression pattern of *Cp2l1* and *Aqp2* expressing cells in kidney of juvenile (P30) *Ae1*<sup>-/-</sup> mouse.** **A-C, G)** Renal cortex. **D-F, H)** Renal inner medulla. Principal cells expressing luminal Aqp2 (aquaporin 2) in renal cortical (**A**) and medullary (**D**) collecting ducts. Nuclear expression pattern of Cp2l1 protein was detected in cortical tubular structures (**B**), while a cell membrane associated and exclusively extranuclear signal for Cp2l1 was detected in inner medullary tubules (**E**). Overlay reveals expression of Cp2l1 in Aqp2 positive and negative cells along the renal collecting ducts (**C, F, G, H**). Nucleic acid stained with DAPI (blue). # Aqp2 negative/Cp2l1 positive cell; arrowhead: Aqp2/Cp2l1 positive cell. **A-F)** Magnification 400x. **G, H)** Magnification 1,000x.

### ***10.1.3 Pendrin is normally expressed in embryonic Ae1 deficient kidney***

Kidney sections of 18 days old Ae1 WT and KO embryos were costained with Aqp2 and pendrin (Fig. 10.19), as well as Aqp2 and the  $\alpha 4$  and B1 subunits of V-ATPase (data not shown). Despite the intriguing abnormal presence of pendrin in the inner medulla of mature (P30 and P84) Ae1 deficient kidneys (Fig. 10.2 and 10.7), pendrin expression in embryonic kidneys (E18) appeared normal (Fig. 10.19). However, the analyzed tubules might be rather derived from the renal cortex, where normal Pds expression was also preserved in Ae1 deficient mature kidneys. Therefore, analysis of the embryonic inner medullary region seems inevitable. However, mRNA expression studies of Ae1 deficient kidneys at different time points of life suggest a late onset (after postnatal day 3, P3) of abnormal expression of acid-base related transporters and signalling molecules in Ae1 KO kidney (Fig. 10.20).



**Fig. 10.19 Expression pattern of Aqp2 and Pds expressing cells in embryonic kidney (E18) of Ae1<sup>-/-</sup> mouse. A)** Principal cells are identified by luminal Aqp2 (aquaporin 2) expression. **B)** Pds (pendrin) expression identifies type B-intercalated cells. **C)** merge. Nucleic acid stained with DAPI (blue). # Aqp2 positive/Pds negative cell; § Aqp2 negative/Pds positive cell. Magnification 1,000x.



#### **10.1.4 *Ae1* deficiency causes late dysregulation of transcription factors**

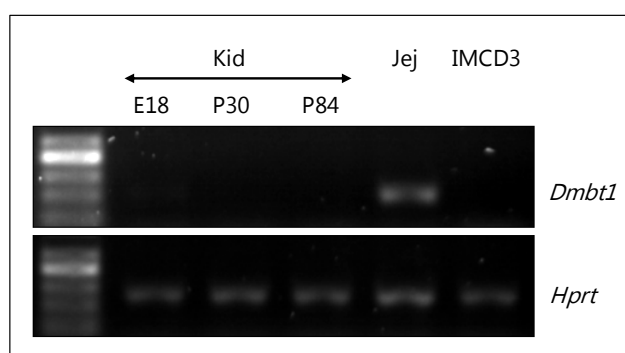
In order to address the question, whether *Ae1* deficiency affects the proper expression of genes regulating CD differentiation, mRNA expression levels of candidate genes were compared at different time points of life (E18, P3, P30, P84) and between *Ae1*<sup>+/+</sup>, *Ae1*<sup>+/-</sup> and *Ae1*<sup>-/-</sup> kidneys. According to chapter 2.7 and 5.2.2, *Foxi1*, *Cp2l1*, *Klf4*, *Bdkrb2*, and *Gdf15* are implicated in terminal differentiation of renal CD epithelium. Furthermore, the renal expression of the acid-base regulatory transporters *Ae1*, *Pds*, *Rhcg* and *B1 subunit of V-ATPase* were investigated because of the known complication of dRTA in *Ae1* deficiency. The results obtained by qRT-RT-PCR are presented in Fig. 10.20. Interestingly, a difference in the mRNA levels of any gene tested between the *Ae1* genotypes was not observed in embryonic kidneys. Moreover, dysregulation of mRNA expression of some candidate genes tested was observed only after postnatal day 3 (P3). As expected and in agreement with the development of dRTA, mRNA of *Pds* and *B1* were significantly downregulated in adult *Ae1* KO kidneys (P84). Surprisingly, there was no difference in mRNA abundance of *Foxi1* or *Cp2l1* in *Ae1* deficient kidneys at any time point of life examined. However, it is of note, that mRNA levels of regulatory molecules involved in p53 mediated signalling pathways, namely *Gdf15*, *Klf4* and *Bdkrb2*, were significantly upregulated in *Ae1* KO kidneys after postnatal day 3 (P3).

**Tab. 10.20 Summary of relative mRNA abundance of genes involved in renal collecting duct differentiation in *Ae1* mutant kidney.** The relative mRNA abundance of several target genes was assessed in male *Ae1* mutant mice at four different time points of life. n= 4-8 animals per genotype and age. Gene expression levels were normalized to *Hprt* expression and calculated in percentage (chapter 9.6). E18: embryonic day 18, P3: postnatal day 3, P30: postnatal day 30, P84: postnatal day 84. Statistical analysis using one way ANOVA and Tukey post-test. \* indicating significant difference compared to *Ae1*<sup>+/+</sup>; # indicating significant difference compared to *Ae1*<sup>+/-</sup>. \*/# p ≤ 0.05, \*\*/## p ≤ 0.01, \*\*\*/### p ≤ 0.001

		% <i>Ae1</i>	% <i>Pds</i>	% <i>Rhcg</i>	% <i>B1</i>	% <i>Foxi</i>	% <i>Cp2l1</i>	% <i>Gdf15</i>	% <i>Klf4</i>	% <i>Bdkrb2</i>
<b>E18</b>	<b><i>Ae1</i><sup>+/+</sup></b>	100.0 ± 33.6	100.0 ± 11.2	100.0 ± 6.8	100.0 ± 14.7	100.0 ± 7.1	100.0 ± 8.8	100.0 ± 16.8	100.0 ± 12.0	100.0 ± 11.6
	<b><i>Ae1</i><sup>+/-</sup></b>	171.9 ± 17.8	94.0 ± 14.3	87.1 ± 11.5	81.0 ± 16.4	104.2 ± 7.4	98.0 ± 11.8	92.5 ± 25.9	100.8 ± 6.9	106.3 ± 28.9
	<b><i>Ae1</i><sup>-/-</sup></b>	1.4 ± 0.5 * ##	95.9 ± 14.2	93.2 ± 7.1	78.6 ± 14.2	114.8 ± 8.7	101.6 ± 11.8	112.3 ± 17.2	109.7 ± 6.8	157.7 ± 26.3
<b>P3</b>	<b><i>Ae1</i><sup>+/+</sup></b>	100.0 ± 23.6	100.0 ± 40.6	100.0 ± 18.6	100.0 ± 6.8	100.0 ± 5.3	100.0 ± 8.1	100.0 ± 12.3	100.0 ± 3.7	100.0 ± 23.1
	<b><i>Ae1</i><sup>+/-</sup></b>	54.6 ± 17.3	70.9 ± 10.5	125.2 ± 9.4	97.6 ± 6.7	98.7 ± 4.7	110.0 ± 6.1	107.7 ± 9.2	98.4 ± 8.9	140.8 ± 14.0
	<b><i>Ae1</i><sup>-/-</sup></b>	0.1 ± 0.0 *	28.1 ± 5.7	78.9 ± 20.9	90.2 ± 8.9	77.5 ± 13.5	82.0 ± 12.8	119.7 ± 18.6	144.5 ± 18.5 #	169.4 ± 24.5
<b>P30</b>	<b><i>Ae1</i><sup>+/+</sup></b>	100.0 ± 22.6	100.0 ± 15.7	100.0 ± 23.2	100.0 ± 14.5	100.0 ± 23.0	100.0 ± 27.5	100.0 ± 9.5	100.0 ± 15.6	100.0 ± 24.4
	<b><i>Ae1</i><sup>+/-</sup></b>	25.8 ± 1.7 **	50.6 ± 3.7 **	62.9 ± 3.1	71.5 ± 7.6	94.0 ± 12.8	74.6 ± 9.6	85.9 ± 7.2	83.5 ± 8.2	92.4 ± 24.4
	<b><i>Ae1</i><sup>-/-</sup></b>	0.3 ± 0.3 ***	78.7 ± 4.3	100.2 ± 15.9	110.8 ± 13.2	164.1 ± 54.5	123.8 ± 26.3	165.9 ± 23.3 * ##	270.1 ± 73.7 * #	309.2 ± 73.0 * #
<b>P84</b>	<b><i>Ae1</i><sup>+/+</sup></b>	100.0 ± 16.6	100.0 ± 12.5	100.0 ± 8.9	100.0 ± 10.4	100.0 ± 13.3	100.0 ± 3.9	100.0 ± 11.7	100.0 ± 6.1	100.0 ± 11.9
	<b><i>Ae1</i><sup>+/-</sup></b>	53.3 ± 8.5 *	127.8 ± 6.7	134.6 ± 4.1	105.9 ± 3.2	111.6 ± 5.3	106.9 ± 8.3	76.9 ± 4.8	86.1 ± 3.6	87.1 ± 13.7
	<b><i>Ae1</i><sup>-/-</sup></b>	0.6 ± 0.4 *** #	72.5 ± 3.5 ##	99.7 ± 20.4	71.5 ± 5.4 #	121.6 ± 9.7	112.1 ± 15.2	201.2 ± 17.6 *** ###	214.7 ± 25.7 *** ###	200.3 ± 54.7 #



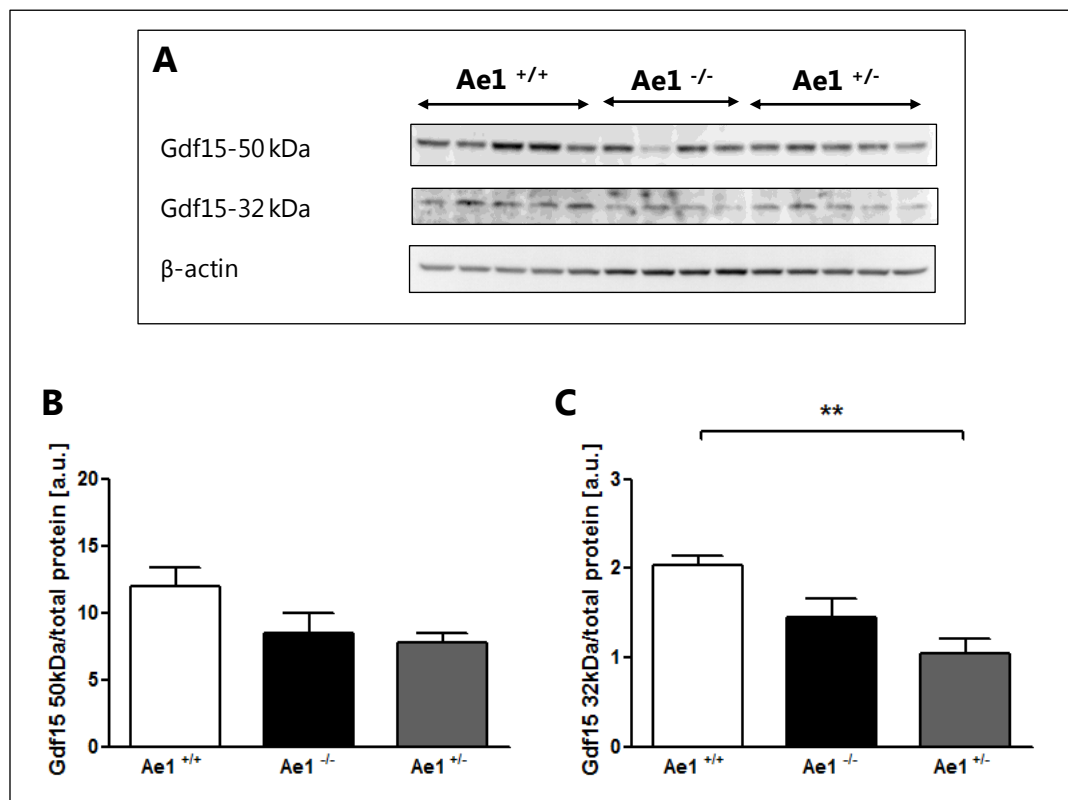
The extra-cellular matrix protein *hensin* (*Dmbt1*) was reported to be crucial for renal intercalated cell differentiation and interconversion (chapter 2.7.5). Since we observed an abnormal expression of Pds in the inner medulla of *Ae1* deficient mice, we aimed to investigate mRNA expression levels of *hensin*. Surprisingly, we could not detect *hensin* mRNA in the mouse kidney at any time point of life (Fig. 10.21). Additionally, *hensin* was not detected in cDNA of the immortalized mouse inner medullary collecting duct cell line IMCD3. However, the primer pair used to detect *hensin* produced the expected product with cDNA obtained from mouse jejunum, confirming specificity and feasibility of the assay. Similar results were obtained using the Taqman gene expression assay from Applied Biosystems for mouse *Dmbt1* (Mm00455996\_1m\*) (data not shown). In order to test, whether mouse strain specific differences may lead to the unexpected absence of *hensin* in C57/Bl6 mouse kidney, total mRNA was extracted from 129SvEv wild-type mouse kidney and the cDNA synthesized was tested (data not shown). Again, we could not detect a RT-RT-PCR primer product for *hensin*. Since a lack of detectable mRNA certainly bears the absence of a related protein product, we have to conclude that *hensin* is not expressed in the kidneys of the mouse strains tested. This is in contrast to the recent report of [99], which was discussed in more detail in chapter 2.7.5.



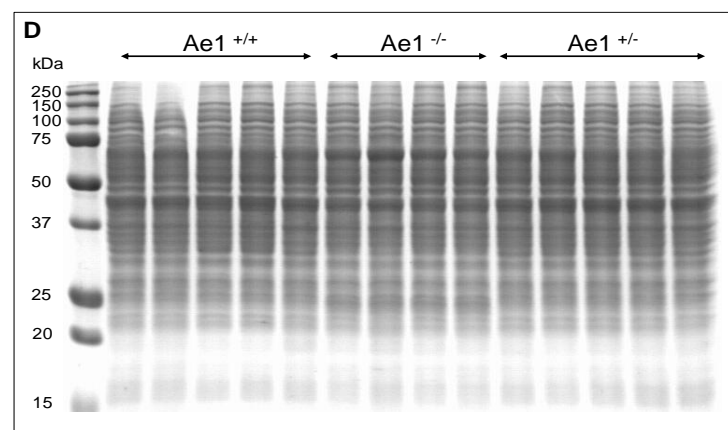
**Fig. 10.21 Lack of detectable *hensin* (*Dmbt1*) mRNA in mouse kidney. Top** qRT-PCR using *hensin* specific primer was performed on kidney cDNA from embryonic (E18), juvenile (P30) and adult (P84) mice, and cDNA obtained from mouse inner medullary collecting duct cell line (IMCD3). The PCR

product was loaded on an Agarosegel and visualized under UV-light. *Hensin* is reported to be highly expressed in the intestine and consistently the expected PCR product was obtained in cDNA of mouse jejunum (Jej). **Bottom** The housekeeping gene *Hprt* was used for additional control.

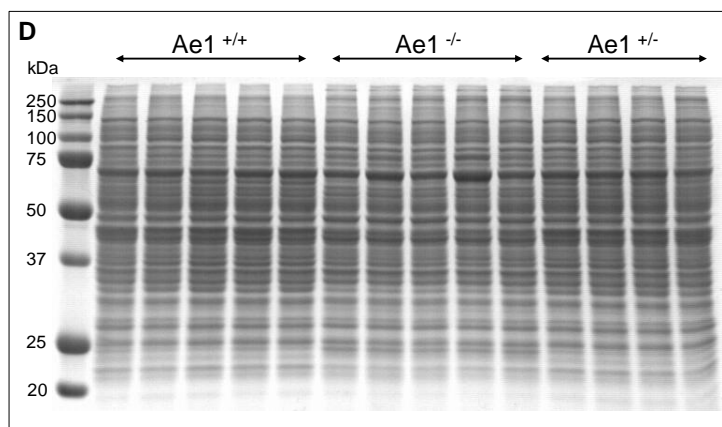
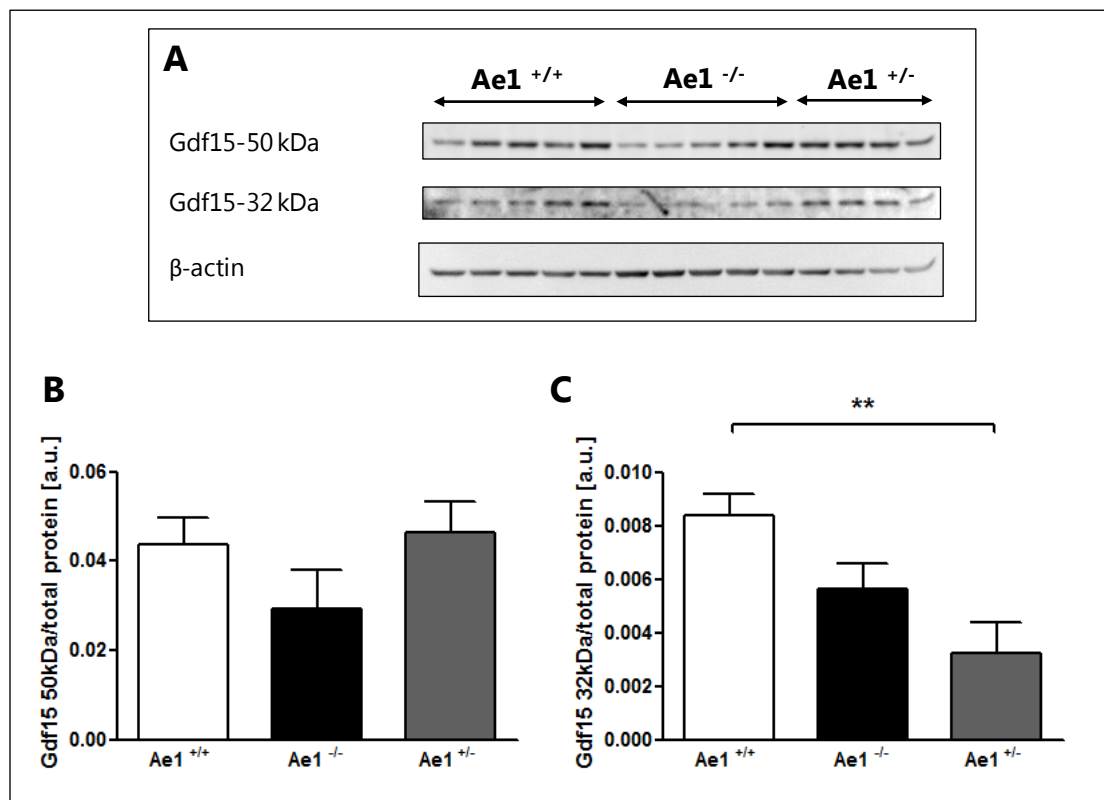
Since *Gdf15* mRNA was significantly upregulated in mature Ae1 KO mice (P30 and P84) compared to WT, the protein abundance was assessed by immunoblotting (Fig. 10.22, 10.23). As illustrated in Fig. 5.2, the Gdf15 precursor peptide undergoes extensive cellular processing, resulting in the biologically active and secreted mature Gdf15 dimer, which is reported to migrate at approximately 30 kDa in electrophoretic SDS-PAGE [181, 225]. Immunoblotting for Gdf15 on cytosolic fractions of P30 and P84 kidneys revealed significant downregulation of the secreted mature Gdf15 dimer, in Ae1<sup>+/-</sup> kidney compared to Ae1<sup>+/+</sup> (Fig. 10.22C, 10.23C), while a probably immature propeptide was not altered [225]. However, the mild decrease in Gdf15 protein abundance in Ae1<sup>-/-</sup> kidneys did not reach significant difference. Cytosolic fractions of mouse liver and placenta homogenates, respectively, were included for positive control (data not shown). Interestingly, the upper 50 kDa band of Gdf15 was exclusively present in the liver, while the lower 32 kDa band of Gdf15 was only detected in placenta samples (data not shown). Thus, a further proof of the molecular identity of the detected Gdf15 related bands in kidney derived samples is requested. In any case, the quality of the Gdf15 immunoblots is not satisfying and needs to be improved.



**Fig. 10.22 Protein abundance of Gdf15 in P84 Ae1 mutant kidneys.**



Representative western blot (**A**) for Gdf15 (growth differentiation factor 15) in cytosolic fractions of P84  $Ae1^{+/+}$ ,  $Ae1^{-/-}$  and  $Ae1^{+/-}$  kidneys. Two Gdf15 related bands were detected, at 50 kDa and 32 kDa, respectively, probably corresponding to immature propeptide and mature dimer. The relative protein abundance of  $\beta$ -actin was significantly higher in  $Ae1^{-/-}$  kidneys, despite loading of equal protein amounts and was therefore not considered for normalization. **B, C**) Densitometry of respective Gdf15 signal intensities, normalized to Coomassie Blue staining of duplicate SDS-PAGE gel (**D**). Statistic analysis using one way ANOVA and Tukey post-test. \*  $p \leq 0.05$ , \*\*  $p \leq 0.01$ , \*\*\*  $p \leq 0.001$ .



**Fig. 10.23 Protein abundance of Gdf15 in P30 Ae1 mutant kidneys.**

Representative western blot (**A**) for Gdf15 (growth differentiation factor 15) in cytosolic fractions of P30 Ae1<sup>+/+</sup>, Ae1<sup>-/-</sup> and Ae1<sup>+/-</sup> kidneys. Two Gdf15 related bands were detected, at 50 kDa and 32 kDa, respectively,

probably corresponding to immature propeptide and mature dimer. The relative protein abundance of β-actin was significantly higher in Ae1<sup>-/-</sup> kidneys, despite loading of equal protein amounts and was therefore not considered for normalization. **B, C**) Densitometry of respective Gdf15 signal intensities, normalized to Coomassie Blue staining of duplicate SDS-PAGE gel (**D**). Statistic analysis using one way ANOVA and Tukey post-test. \*  $p \leq 0.05$ , \*\*  $p \leq 0.01$ , \*\*\*  $p \leq 0.001$ .

## ***10.2 Renal adaptation to chronic metabolic acidosis***

### ***10.2.1 Metabolic parameters***

As described in chapter 5.2, acid-base disturbances are known to induce several adaptive responses in the kidney, including a cellular remodelling of the CD epithelium. Therefore, a possible impact on induced chronic metabolic acidosis, lasting for 8 weeks, was investigated and finally compared with observations in Ae1 deficiency. Metabolic parameters of experimental mice (P84) are summarized in Tab. 10.24 and confirmed the expected adaptation to metabolic acidosis induced by addition of  $\text{NH}_4\text{Cl}$  to the food. The health status of the animals was tightly monitored during the experimental procedure by subjective observations (appearance of fur, eyes and activity). Furthermore, adolescence of the acidotic ( $\text{NH}_4\text{Cl}$ ) animals seemed normal, since the bodyweight was unaffected by the unusual diet. The daily water intake and respective urinary output was increased in acidotic animals as previously reported [219]. The increased blood plasma chloride levels as well as the increased secretion of urinary  $\text{NH}_3/\text{NH}_4^+$  reflected the chronic oral uptake of  $\text{NH}_4\text{Cl}$  with the food in acidotic mice. The increased renal secretion of acid equivalents to the urine, indicated by the lowered urinary pH and increased urinary  $\text{NH}_3/\text{NH}_4^+$  excretion, completely compensated the induced metabolic acidosis. Therefore, blood pH,  $\text{pCO}_2$  and  $\text{HCO}_3^-$  concentrations, as well as the Anion Gap were normal in  $\text{NH}_4\text{Cl}$  loaded mice compared to Control.

**Tab. 10.24 Metabolic and urine parameters of Control and 8 weeks NH<sub>4</sub>Cl loaded C57/Bl6 (P84)****mice.** Statistical analysis using Student's *t*-test (n = 10 per group).\*  $p \leq 0.05$ , \*\*  $p \leq 0.01$ , \*\*\*  $p \leq 0.001$ .

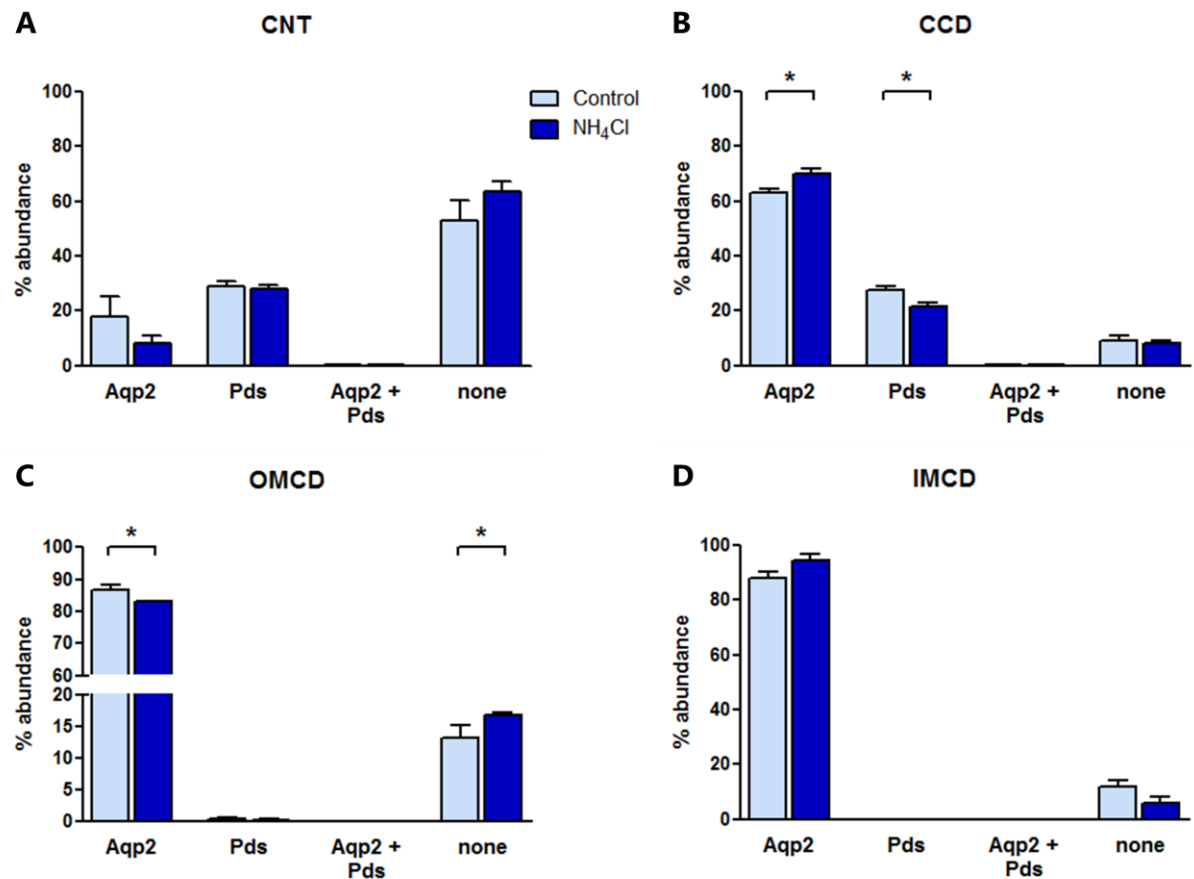
	<b>Control</b>	<b>NH<sub>4</sub>Cl</b>
Bodyweight (g)	26.16 ± 0.55	25.10 ± 0.36
H <sub>2</sub> O Intake (g/g BW)	0.18 ± 0.01	0.26 ± 0.01 ***
Food Intake (g/g BW)	0.17 ± 0.01	0.16 ± 0.01
Stool (g/g BW)	0.04 ± 0.00	0.04 ± 0.00
Urine (g/g BW)	0.04 ± 0.01	0.09 ± 0.01 ***
<b>Blood</b>		
pH	7.35 ± 0.02	7.39 ± 0.07
pCO <sub>2</sub> (mmHG)	45.86 ± 2.46	45.20 ± 2.91
pO <sub>2</sub> (mmHG)	34.94 ± 0.88	41.63 ± 2.17
HCO <sub>3</sub> <sup>-</sup> (mM)	24.52 ± 0.71	22.96 ± 1.54
Na <sup>+</sup> (mM)	144.60 ± 0.98	146.20 ± 0.37
K <sup>+</sup> (mM)	4.02 ± 0.21	3.98 ± 0.07
Cl <sup>-</sup> (mM)	107.00 ± 0.84	110.60 ± 1.25 *
Ca <sup>++</sup> (mM)	1.15 ± 0.08	1.04 ± 0.06
Hematocrit (%)	45.26 ± 0.40	47.12 ± 1.61
Anion Gap	13.16 ± 1.41	12.60 ± 0.30
<b>Urine</b>		
Creatinine (mg/dl)	32.08 ± 2.38	16.38 ± 1.08 ***
pH	5.83 ± 0.02	5.46 ± 0.01 ***
HCO <sub>3</sub> <sup>-</sup> (mM)/Crea (mg/dl)	0.008 ± 0.0009	0.005 ± 0.0005 *
NH <sub>3</sub> /NH <sub>4</sub> <sup>+</sup> (mM)/Crea (mg/dl)	2.05 ± 0.14	26.98 ± 0.63 ***

### ***10.2.2 Renal expression pattern of acid-base transporters during chronic metabolic acidosis***

The renal expression pattern of Pds, V-ATPase (represented by a4 and B1 subunit) and Ae1 was assessed by immunofluorescence and the relative abundance of individually labelled cells of Control and NH<sub>4</sub>Cl treated mice were compared (Fig. 10.25 to 10.28). Furthermore, the relative protein abundance of the afore-mentioned transporters, as well as Gdf15 and Foxi1, was determined by immunoblotting (Fig. 10.29 to 10.34). Finally, the relative mRNA abundance of candidate genes expressed in kidneys was compared between Control and 8 weeks NH<sub>4</sub>Cl treated mice and Ae1 mutant mice, respectively (Fig. 10.35, 10.36).

The obtained results clearly show the renal compensation of the chronic oral acid load in treated C57/Bl6 mice. Moreover, the adaptive responses of the kidney to chronic metabolic acidosis are significantly different to the observations made in Ae1 deficiency.

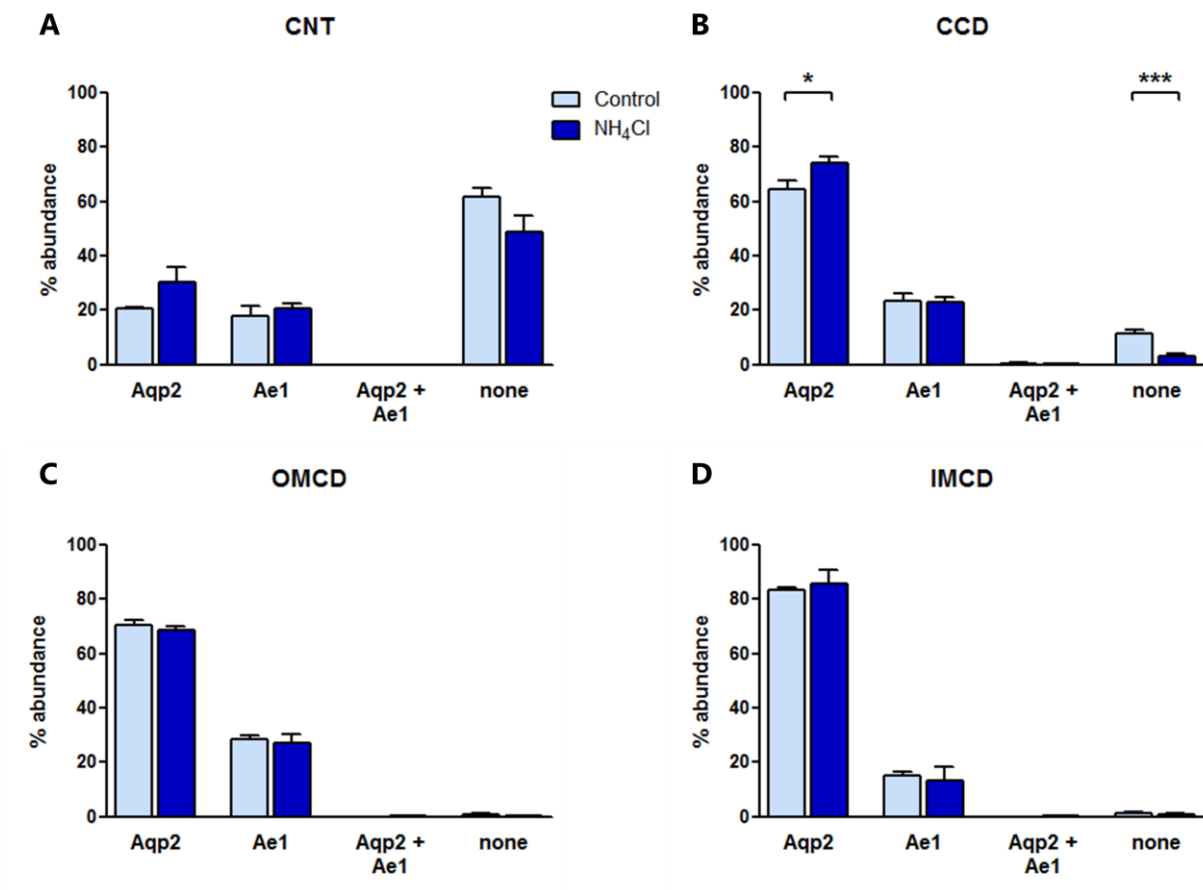
As expected and shown in Fig. 10.25, the relative abundance of Pds expressing type B-IC decreased significantly upon chronic oral acid loading. However, this adaptation was totally accounted by alterations of the relative cell number in the renal CCD, since the abundance of Pds expressing type B-ICs was not changed in the renal CNT, OMCD or IMCD of NH<sub>4</sub>Cl loaded mice compared to Control. The expected decrease of type B-ICs in the renal CCD of acidotic animals was accompanied by an increase of Aqp2 labelled principal cells (Fig. 10.25B). In the renal OMCD, the relative abundance of Aqp2<sup>+</sup> cells was reduced in acidotic kidneys, while unlabelled (none) cells increased concomitantly (Fig. 10.25C). A similar, but not yet significantly different trend was observed in renal IMCD of acid loaded mice (Fig. 10.25D). As expected and importantly in contrast to Ae1 deficient mice, Pds expressing cells were not detected at all in the IMCD of chronic NH<sub>4</sub>Cl treated animals (Fig. 10.3D, 10.8D, 10.25D).



**Fig. 10.25 Relative distribution of Aqp2 and Pds expressing cells in kidneys of Control and 8 weeks NH<sub>4</sub>Cl treated mice (P84).** Summary of cell counts assessing the relative abundance of Aqp2 (aquaporin 2) and Pds (pendrin) positive cells in the renal CNT, CCD, OMCD and IMCD of Control and NH<sub>4</sub>Cl treated mice. In total, 7,882 cells were counted in Control kidneys (n=5) and 4,014 cells in NH<sub>4</sub>Cl treated kidneys (n=5). Statistical analysis using Student's *t*-test. \*  $p \leq 0.05$ , \*\*  $p \leq 0.01$ , \*\*\*  $p \leq 0.001$ . CNT: connecting tubule, CCD: cortical collecting duct, OMCD: outer medullary collecting duct, IMCD: inner medullary collecting duct



Interestingly, the relative abundance of Ae1 positive (Ae1<sup>+</sup>) cells remained constant upon chronic NH<sub>4</sub>Cl loading (Fig. 10.26). However, similar to the observations made in the Pds staining series (Fig. 10.25B), a significant increase in Aqp2<sup>+</sup> cells on the expense of unlabelled (none) cells was present in renal CCD of acid loaded mice (Fig. 10.26B). The unlabelled (none) cells of the Ae1 staining series comprise all types of ICs with the exception of type A-ICs as defined by the respective marker proteins. Therefore, the data suggest that principal cells in the renal CCD increase on the expense of type B-ICs upon chronic acid stimuli. Immunologic detection of the V-ATPase, a more general marker of renal ICs, confirmed the conclusion.

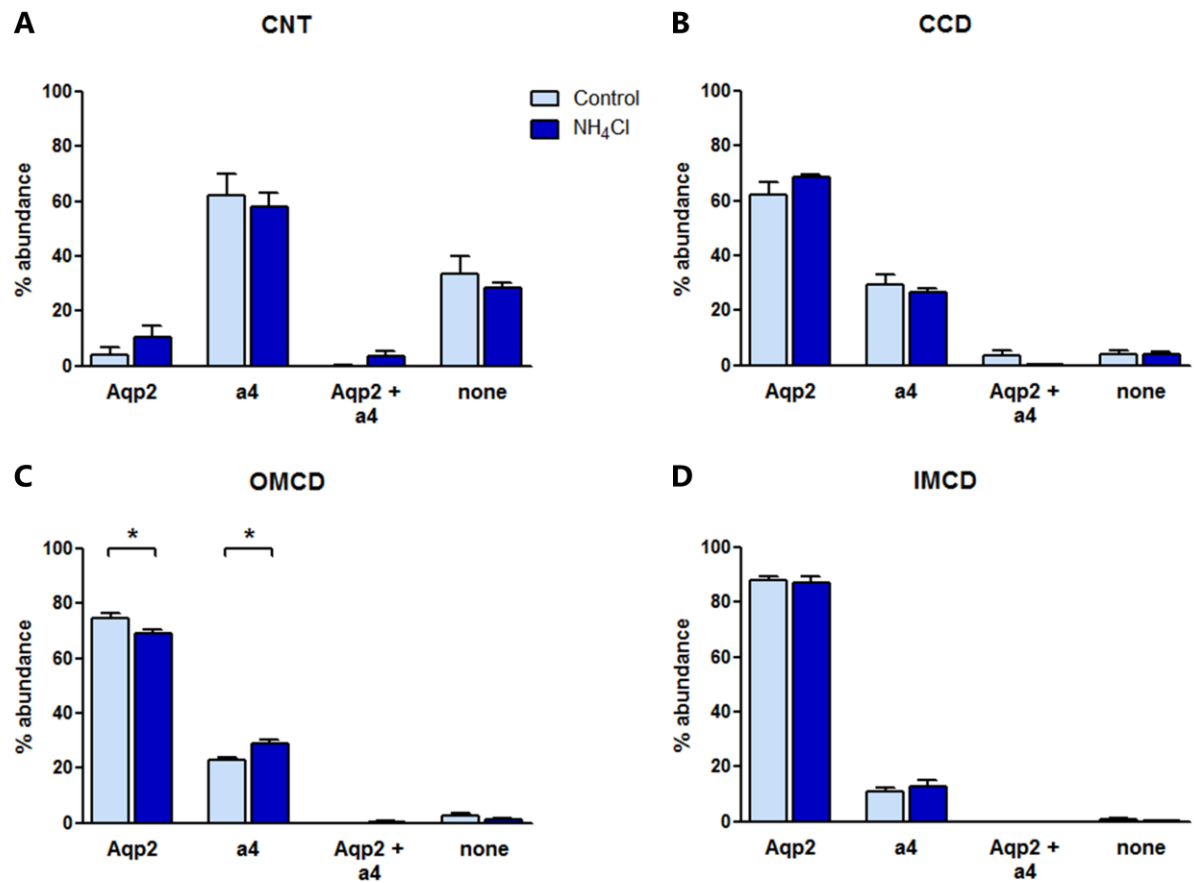


**Fig. 10.26 Relative distribution of Aqp2 and Ae1 expressing cells in kidneys of Control and 8 weeks NH<sub>4</sub>Cl treated mice (P84).** Summary of cell counts assessing the relative abundance of Aqp2 (aquaporin 2) and Ae1 (anion exchanger 1) positive cells in the renal CNT, CCD, OMCD and IMCD of Control and NH<sub>4</sub>Cl treated mice. In total, 3,926 cells were counted in Control kidneys (n=5) and 2,913 cells in NH<sub>4</sub>Cl treated kidneys (n=5). Statistical analysis using Student's *t*-test. \*  $p \leq 0.05$ , \*\*  $p \leq 0.01$ , \*\*\*  $p \leq 0.001$ .

CNT: connecting tubule, CCD: cortical collecting duct, OMCD: outer medullary collecting duct, IMCD: inner medullary collecting duct

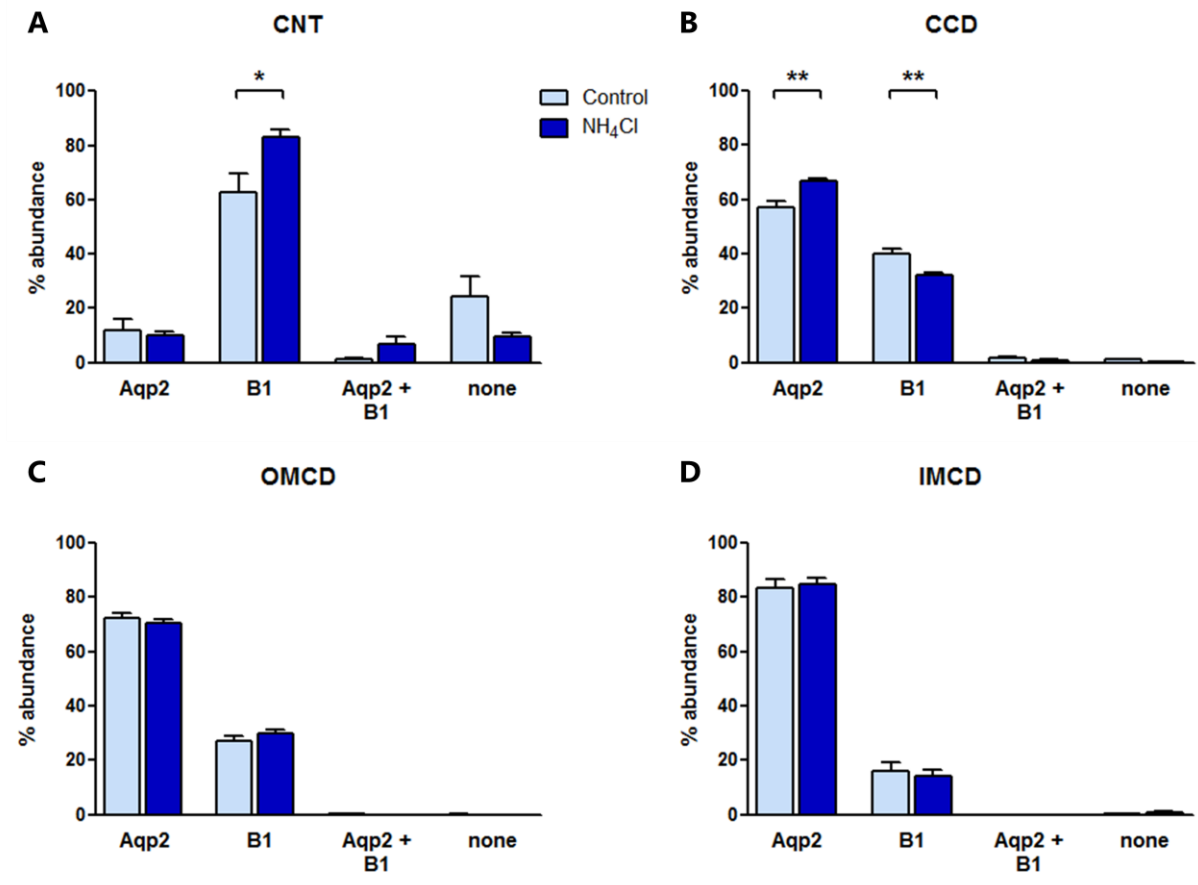
The V-ATPase is composed of distinct subunits (Fig. 5.1) and ubiquitously expressed in different tissues. The  $\alpha 4$  subunit of the V-ATPase is specifically expressed in the kidney, while the B1 subunit is only present in renal ICs of the CD system, thus representing a more specific molecular marker. The cellular localization of the V-ATPase differs among renal ICs, matching their function in acid-base regulation. However, the subcellular localization of the V-ATPase within the renal ICs was not further included in the quantitative analysis of the immunologic staining series. Here, cells positive for the  $\alpha 4$  ( $\alpha 4^+$ ) or B1 ( $B1^+$ ) subunit combine all manifestations of the cellular V-ATPase localization (i.e. apical, cytosolic, basolateral). Comparing the relative abundance of V-ATPase labelled cells in acidotic kidneys, it is of note, that the  $\alpha 4$  and B1 subunit of V-ATPase are differentially regulated in distinct parts of the distal tubule (Fig. 10.27, 10.28). This phenomenon was already described during nephrogenesis by Jouret *et al* [29]. The relative abundance of cells positive for the  $\alpha 4$  subunit of V-ATPase increased in the renal OMCD of  $\text{NH}_4\text{Cl}$  treated mice, while  $\text{Aqp2}^+$  decreased (Fig. 10.27C). However, there were no significant differences observed between Control and acidotic kidneys regarding the  $\alpha 4^+$  or  $\text{Aqp2}^+$  cell abundance in other distal tubule segments (Fig. 10.27A, B, D). In contrast, the relative abundance of  $B1^+$  cells increased significantly in the renal CNT of acidotic mice, maybe to some extent on the expense of unlabelled (none) cells although the reduction did not reach statistical relevance (Fig. 10.28A). However, the renal CNT of mice is a segment, which consists of a variety of distinct and transitory cell types. Hence, it is likely that some less defined cell type expresses the B1 subunit of V-ATPase upon chronic metabolic acidosis. Consistent with observations of the Pds staining series (Fig. 10.25), the relative abundance of  $B1^+$  cells of acidotic CCD was decreased and accompanied by a respective increase in  $\text{Aqp2}^+$  cells (Fig. 10.28B). This observation confirms the previous conclusion that principal cells of the renal CCD increase in abundance, on the expense of type B-ICs upon chronic acid loading. Apart from that, the relative abundance of  $B1^+$  cells in OMCD and IMCD was similar in Control and  $\text{NH}_4\text{Cl}$  treated animals (Fig. 10.28C, D).

Notably and in contrast to Ae1 deficiency, Pds expressing cells remained absent in chronic metabolic acidosis. Furthermore, an interconversion of type A-ICs from type B-ICs upon chronic metabolic acidosis is not likely, as the relative abundance of type A-ICs did not change upon prolonged acid load. Instead of, an adaptive increase in PCs of the renal CCD was observed with a concomitant decrease in type B-ICs. However, whether this is a result of proliferation and apoptosis of the respective cell types or from cell type interconversion remains speculative.



**Fig. 10.27 Relative distribution of Aqp2 and a4 expressing cells in kidneys of Control and 8 weeks NH<sub>4</sub>Cl treated mice (P84).** Summary of cell counts assessing the relative abundance of Aqp2 (aquaporin 2) and a4 (a4 subunit of V-ATPase) positive cells in the renal CNT, CCD, OMCD and IMCD of Control and NH<sub>4</sub>Cl treated mice. In total, 5,482 cells were counted in Control kidneys (n=5) and 4,822 cells in NH<sub>4</sub>Cl treated kidneys (n=5). Statistical analysis using Student's *t*-test. \* *p* ≤ 0.05, \*\* *p* ≤ 0.01, \*\*\* *p* ≤ 0.001.

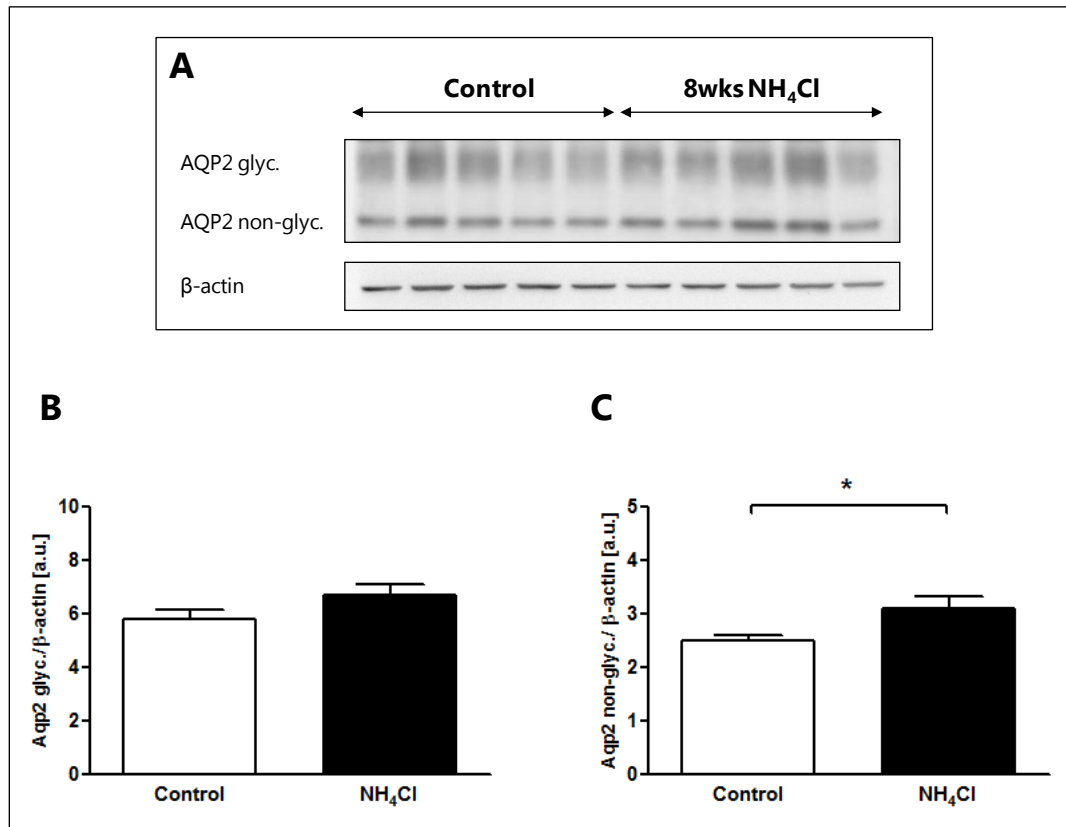
CNT: connecting tubule, CCD: cortical collecting duct, OMCD: outer medullary collecting duct, IMCD: inner medullary collecting duct



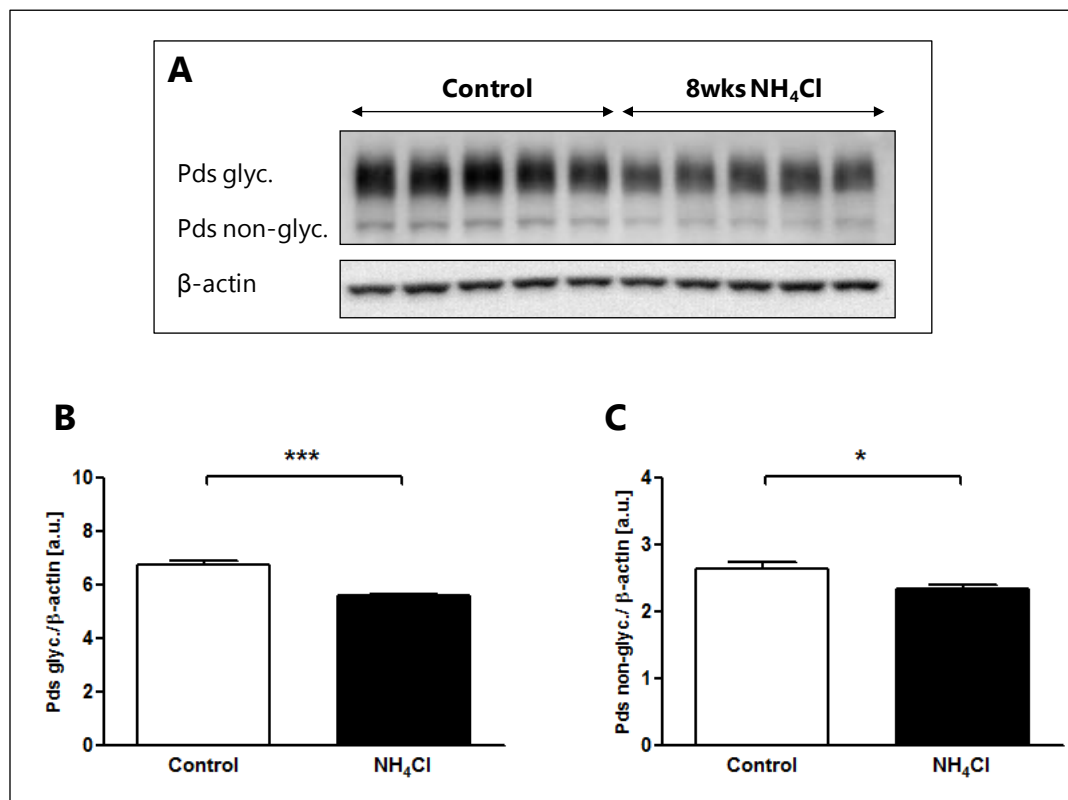
**Fig. 10.28 Relative distribution of Aqp2 and B1 expressing cells in kidneys of Control and 8 weeks NH<sub>4</sub>Cl treated mice (P84).** Summary of cell counts assessing the relative abundance of Aqp2 (aquaporin 2) and B1 (B1 subunit of V-ATPase) positive cells in the renal CNT, CCD, OMCD and IMCD of Control and NH<sub>4</sub>Cl treated mice. In total, 4,714 cells were counted in Control kidneys (n=5) and 4,543 cells in NH<sub>4</sub>Cl treated kidneys (n=5). Statistical analysis using Student's *t*-test. \*  $p \leq 0.05$ , \*\*  $p \leq 0.01$ , \*\*\*  $p \leq 0.001$ .

CNT: connecting tubule, CCD: cortical collecting duct, OMCD: outer medullary collecting duct, IMCD: inner medullary collecting duct

Immunoblotting of total kidney membrane preparations confirmed the results obtained by immunofluorescent labelling. Again, Aqp2 protein abundance was increased in acidotic kidneys (Fig. 10.29), while Pds abundance was decreased (Fig. 10.30). Moreover, the relative protein abundance of Ae1 and the B1 subunit of V-ATPase did not change upon chronic metabolic acidosis (Fig. 10.31, 10.32).

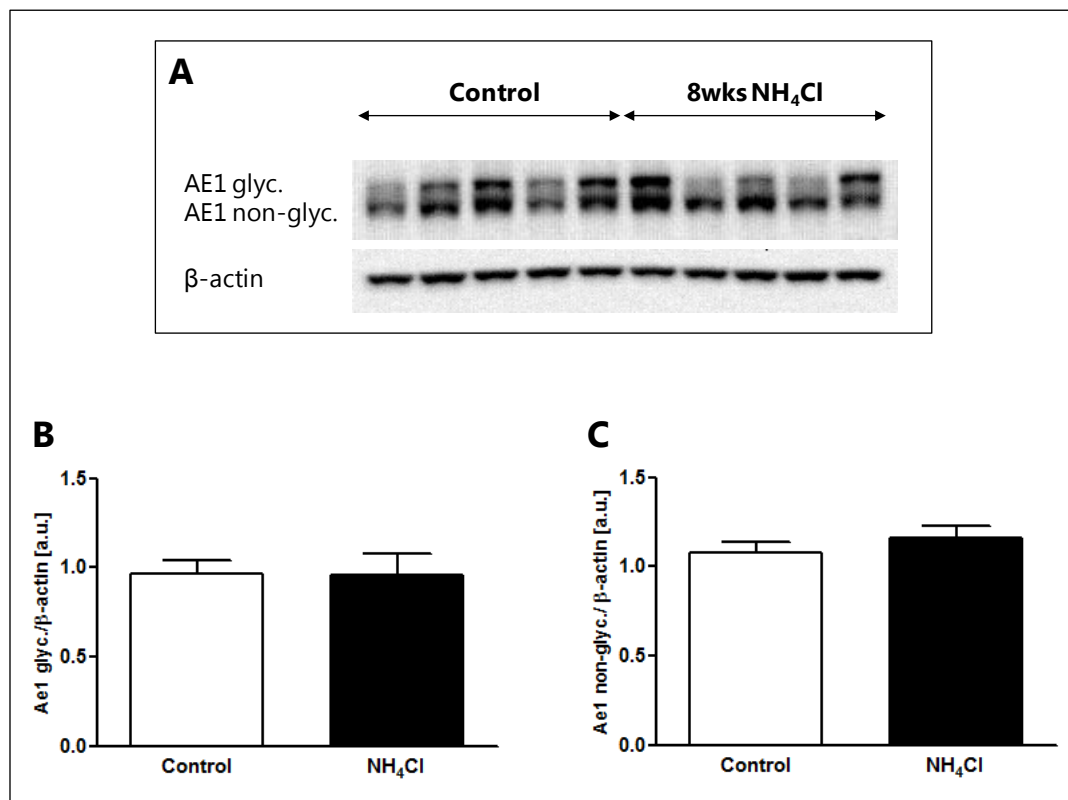


**Fig. 10.29 Relative protein abundance of Aqp2 in kidneys of Control and 8 weeks NH<sub>4</sub>Cl treated mice (P84).** **A**) Representative immunoblot of Aqp2 (aquaporin 2) on total kidney membrane preparation from Control and 8 weeks NH<sub>4</sub>Cl treated mice, β-actin served as loading control. **B, C**) Densitometry. Statistical analysis using Student's *t*-test (*n* = 5 per group). \* *p* ≤ 0.05, \*\* *p* ≤ 0.01, \*\*\* *p* ≤ 0.001.

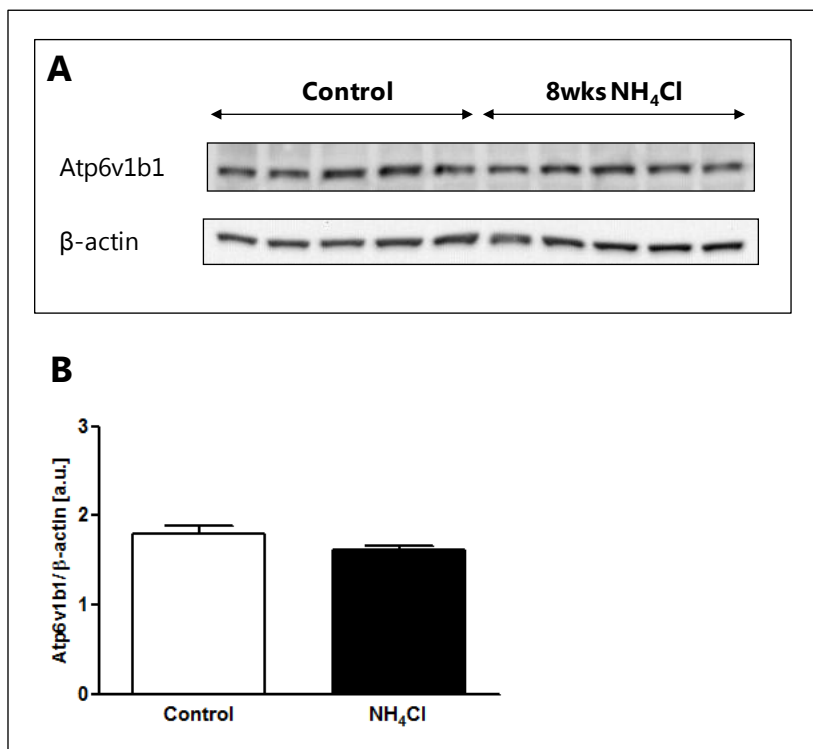


**Fig. 10.30 Relative protein abundance of Pds in kidneys of Control and 8 weeks NH<sub>4</sub>Cl treated mice (P84).** **A**) Representative immunoblot of Pds (pendrin) on total kidney membrane preparation from Control and 8 weeks NH<sub>4</sub>Cl treated mice, β-actin served as loading control. **B, C**) Densitometry. Statistical analysis using Student's *t*-test (n = 5 per group). \* *p* ≤ 0.05, \*\* *p* ≤ 0.01, \*\*\* *p* ≤ 0.001.



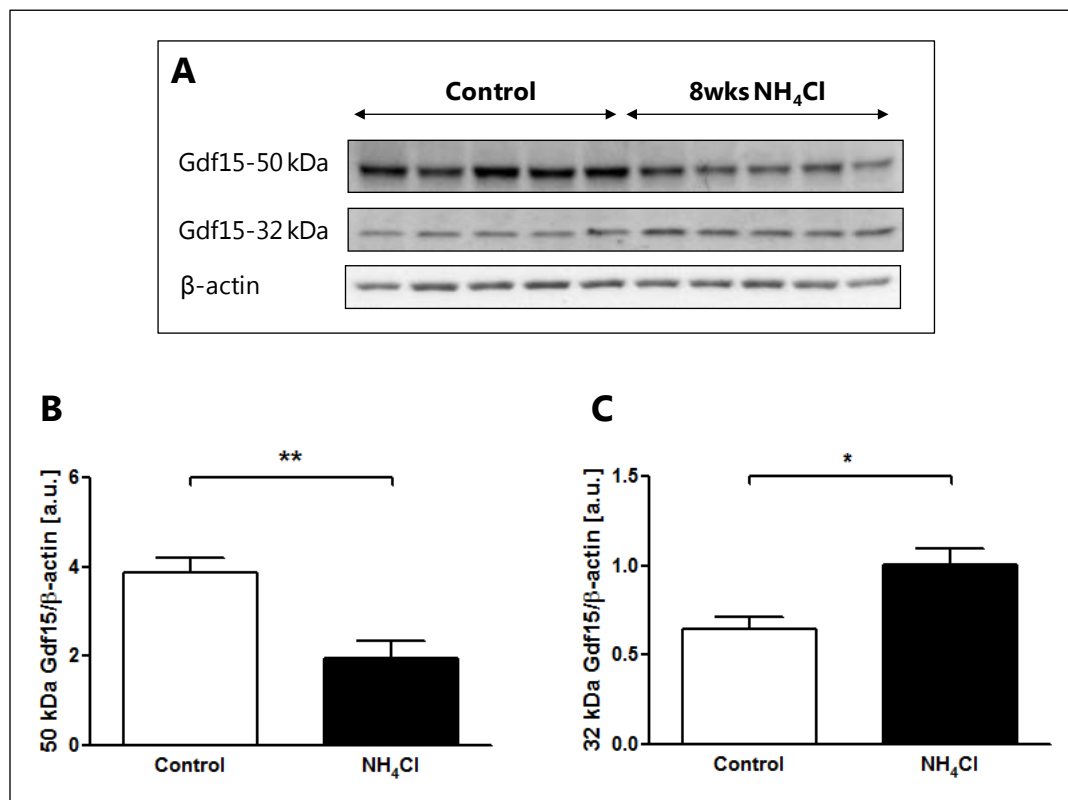


**Fig. 10.31 Relative protein abundance of Ae1 in kidneys of Control and 8 weeks NH<sub>4</sub>Cl treated mice (P84).** **A**) Representative immunoblot of Ae1 (anion exchanger 1) on total kidney membrane preparation from Control and 8 weeks NH<sub>4</sub>Cl treated mice, β-actin served as loading control. **B, C**) Densitometry. Statistical analysis using Student's *t*-test (n = 5 per group). \* *p* ≤ 0.05, \*\* *p* ≤ 0.01, \*\*\* *p* ≤ 0.001.



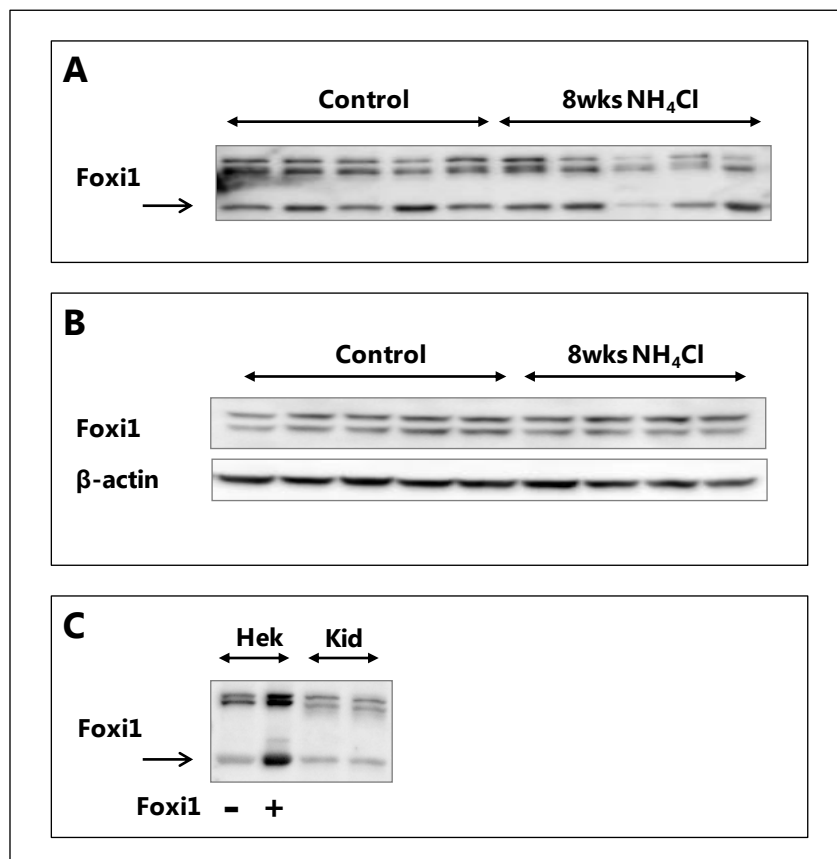
**Fig. 10.32 Relative protein abundance of B1 subunit of V-ATPase in kidneys of Control and 8 weeks NH<sub>4</sub>Cl treated mice (P84).** **A)** Representative immunoblot of B1 subunit of V-ATPase (ATP6v1b1) on total kidney membrane preparation from Control and 8 weeks NH<sub>4</sub>Cl treated mice, β-actin served as loading control. **B)** Densitometry. Statistical analysis using Student's *t*-test (n = 5 per group). \* *p* ≤ 0.05, \*\* *p* ≤ 0.01, \*\*\* *p* ≤ 0.001.

In addition to alterations of acid-base transporters, Gdf15 protein abundance was significantly regulated in chronic metabolic acidosis (Fig. 10.33). A supposedly immature propeptide was detected at 50 kDa and found to be significantly downregulated upon chronic acidosis. Consistently, the probably mature secreted dimer of Gdf15 at 32 kDa was significantly upregulated in acidotic kidneys, while *Gdf15* mRNA was not changed (Fig. 10.35J). However, whether the regulated supposedly mature dimer is really present as a secreted signalling peptide of Gdf15 circulating in the blood is not known, since serum levels of Gdf15 were not assessed in Control or acidotic mice. Furthermore, the function of Gdf15 in adaptive responses of the kidney during chronic metabolic acid-base disturbances is not yet resolved. In the early adaptive response of the kidney to metabolic acidosis, Gdf15 was suggested to be involved in the initiation of proliferation (chapter 5.2.2). However, this function seems unlikely in the case of chronic (8 weeks) acidosis, as the initial increase of the proliferation rate might be already terminated and sustained at low comparable rates of Control kidneys. Moreover, Gdf15 is involved in the regulation of cellular processes upon manifold stress stimuli [181]. Hence, a regulatory role of Gdf15 in the kidney of chronic acidotic mice might be rather involved in processes that reduce cellular stress.



**Fig. 10.33 Relative protein abundance of Gdf15 in kidneys of Control and 8 weeks NH<sub>4</sub>Cl treated mice (P84).** **A**) Representative immunoblot of Gdf15 (growth differentiation factor 15) on cytosolic fractions of total kidney homogenates extracted from Control and 8 weeks NH<sub>4</sub>Cl treated mice, β-actin served as loading control. **B, C**) Densitometry. Statistical analysis using Student's *t*-test (n = 5 per group). \* *p* ≤ 0.05, \*\* *p* ≤ 0.01, \*\*\* *p* ≤ 0.001.

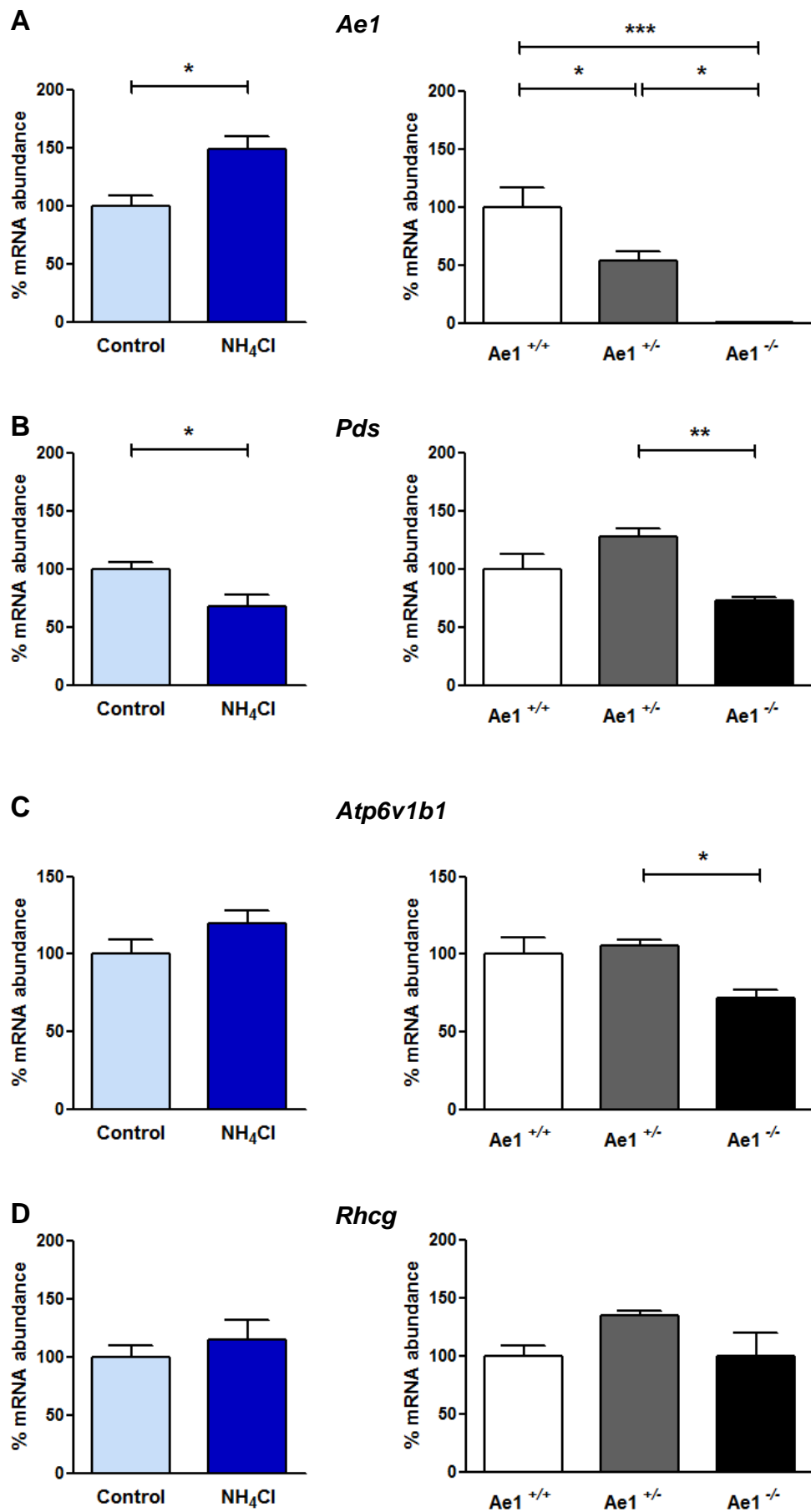
In order to assess the relative protein abundance of Foxi1 in Control and acidotic kidneys, immunoblotting of nuclear protein extracts and cytosolic fractions from kidney homogenates were performed. Transiently *Foxi1* transfected Hek293T cells served as a positive control for the pattern of Foxi1 protein and derivatives in electrophoretic SDS-PAGE (Fig. 10.34C). Here, three Foxi1 related bands were detected by western blot, of which one matched the expected molecular weight of Foxi1 (41 kDa). The other two Foxi1 related bands were detected at higher molecular weights and are of unknown origin. However, in kidney nuclear extracts all three bands were detected and were found expressed at comparable levels in Control and NH<sub>4</sub>Cl treated animals (Fig. 10.34A). Interestingly, Foxi1 immunoblotting on kidney cytosolic fractions of total homogenate showed a distinct pattern (Fig. 10.34B). The lowest band, corresponding to the expected molecular weight was missing. Anyhow, the two higher Foxi1 related bands of unknown origin were present but not regulated upon acid-base disturbance. Future experiments are necessary to solve the character of the Foxi1 related bands detected by immunoblotting. Foxi1 was shown to be capable of remodelling chromatin structures and to remain bound to condensed chromatin during mitosis [226]. However, inactive or premature forms of Foxi1 might also be present outside the nucleus, as indicated by the detection of Foxi1 related proteins with higher molecular weight in the cytosolic fraction of kidney homogenates.



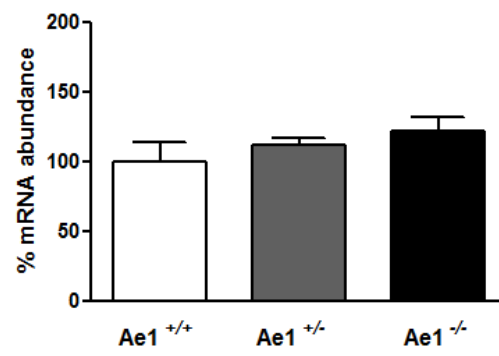
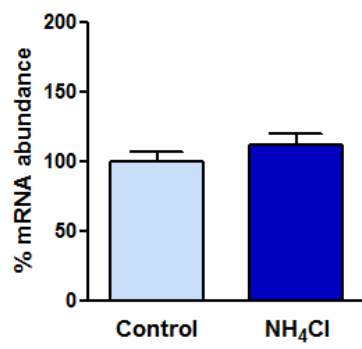
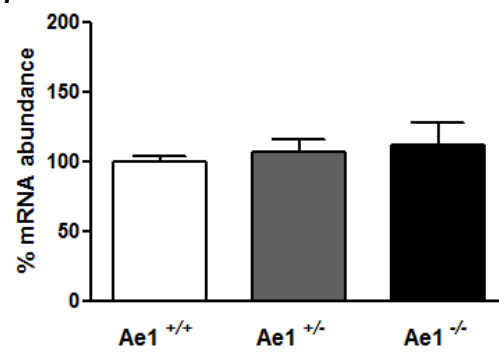
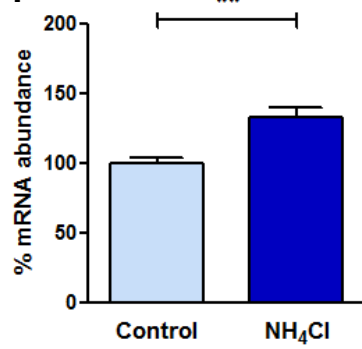
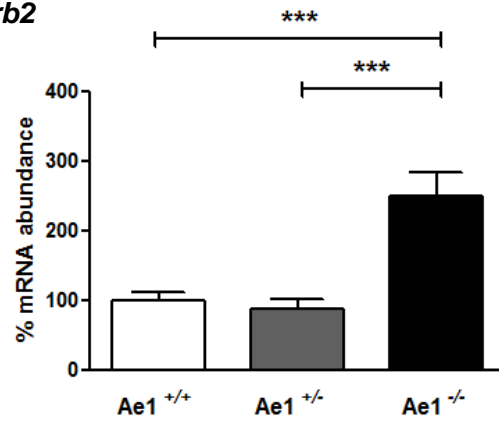
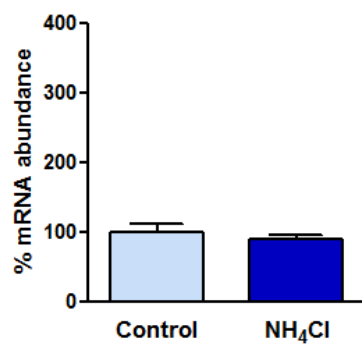
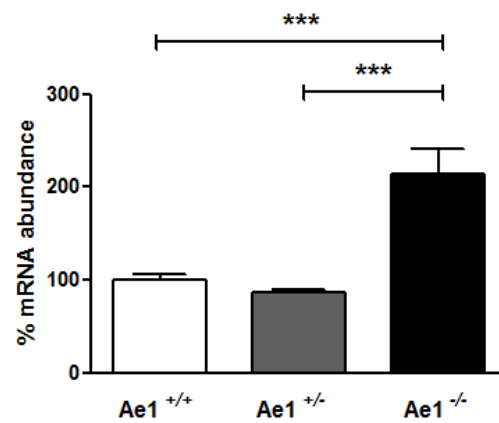
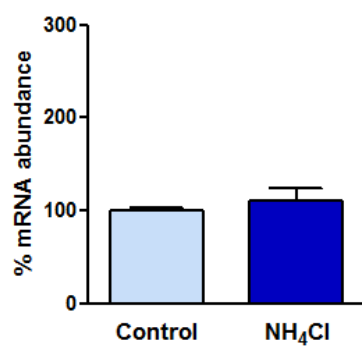
**Fig. 10.34 Relative protein abundance of Foxi1 in kidneys of Control and 8 weeks NH<sub>4</sub>Cl treated mice (P84).** **A-C** Representative immunoblot of Foxi1 (forkheadbox protein 1). **A**) Nuclear protein extracts from total kidneys of Control and 8 weeks NH<sub>4</sub>Cl treated mice. **B**) Cytosolic fractions of total kidney homogenates from Control and 8 weeks NH<sub>4</sub>Cl treated mice with β-actin as loading control. **C**) Nuclear protein extracts from untransfected (Foxi1 -) and transiently transfected Hek293T cells (Foxi1 +) and total kidney (Kid) (P30). arrow: indicates expected molecular weight of Foxi1 (41 kDa), only detectable in nuclear protein extracts.

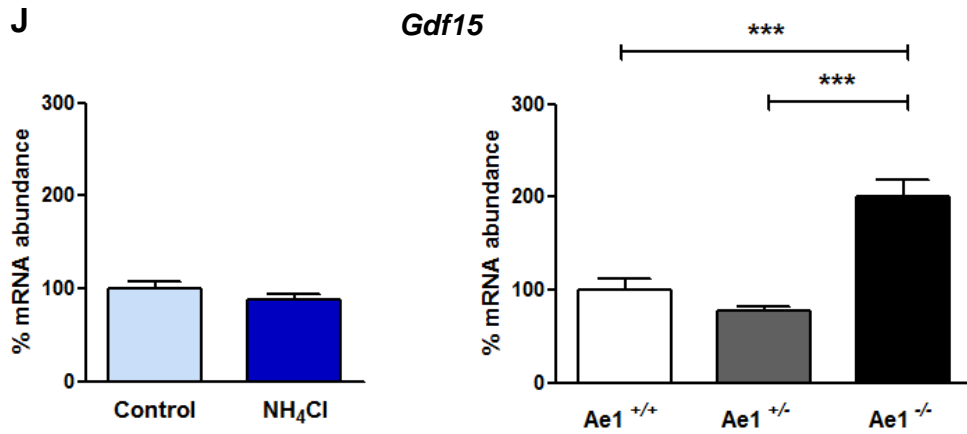
Addressing the question whether chronic acidosis is capable of altering the differentiation of renal IC, total mRNA of Control and NH<sub>4</sub>Cl treated mouse kidney was extracted and qRT-PCR was performed for the above introduced candidate genes of acid-base transporters and transcription factors or signalling molecules. Finally, the results were compared with data obtained from Ae1 mutant mouse kidney. The relative mRNA expression levels are presented in bar graphs of Fig. 10.35A-J and a simplified tabular expression of the results is shown in Fig. 10.36. Consistent with the chronic acidosis, relative mRNA levels of *Ae1* (Fig. 10.35A) were upregulated in NH<sub>4</sub>Cl loaded mice compared to Control, while *Pds* was downregulated (Fig. 10.35B). Similarly, relative mRNA expression of *Pds* was reduced in Ae1 KO mice. The relative abundance of the *B1 subunit of V-ATPase* was not altered in NH<sub>4</sub>Cl loaded animals compared to Control (Fig. 10.35C). However, the *B1 subunit of V-ATPase* was significantly downregulated in Ae1 KO kidneys. Furthermore, *Rhcg* mRNA levels were not changed significantly in chronic metabolic acidosis nor Ae1 related dRTA (Fig. 10.35D). Interestingly, relative *Foxi1* mRNA abundance was not different in NH<sub>4</sub>Cl treated mice compared to Control or Ae1 mutant mice (Fig. 10.35E). Strikingly, relative *Cp2l1* mRNA abundance was upregulated in chronic metabolic acidosis but not in Ae1 KO (Fig. 10.35F). Moreover, relative mRNA levels of *Bdkrb2*, *Klf4* and *Gdf15* were significantly upregulated in kidneys Ae1 KO mice compared to WT, but not in kidneys of NH<sub>4</sub>Cl loaded mice compared to Control (Fig. 10.35G-J).

Hence, the investigated genes are differentially regulated during chronic metabolic acidosis and Ae1 related dRTA as also shown in the simplified table (Fig. 10.36).





**E*****Foxi1*****F*****Cp2l1*****G*****Bdkrb2*****H*****Klf4***



**Fig. 10.35 Relative renal mRNA abundance of target genes during chronic metabolic acidosis and dRTA.** Left bar graph represents mRNA expression levels in C57/Bl6 Control and NH<sub>4</sub>Cl treated mice. Right bar graph shows renal mRNA expression levels of Ae1 mutant mice, respectively. Statistical analysis using Student's *t*-test (n = 4-5 animals per group). \*  $p \leq 0.05$ , \*\*  $p \leq 0.01$ , \*\*\*  $p \leq 0.001$ .

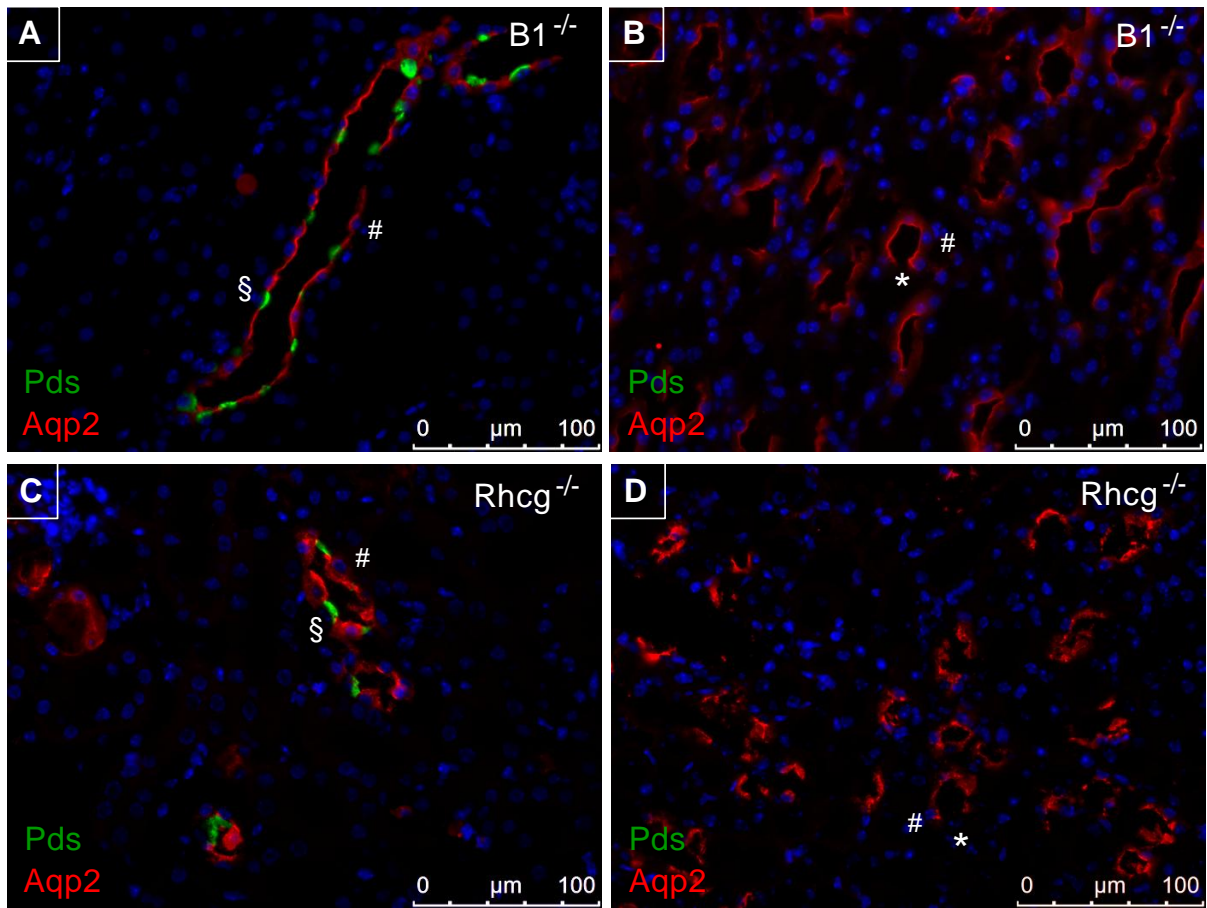
relative change in % mRNA abundance		
gene	Control vs NH <sub>4</sub> Cl	Ae1 WT vs KO
<i>Ae1</i>	↑	↓↓
<i>Pds</i>	↓	↔
<i>Atp6v1b1</i>	↔	↔
<i>Rhcg</i>	↔	↔
<i>Foxi1</i>	↔	↔
<i>Cp2l1</i>	↑↑	↔
<i>Gdf15</i>	↔	↑↑
<i>Klf4</i>	↔	↑↑
<i>Bdkrb2</i>	↔	↔

**Fig. 10.36 Summarized results from Fig. 10.35** representing relative changes in mRNA expression levels of NH<sub>4</sub>Cl treated mice relative to C57/Bl6 Control and Ae1 KO mice relative to Ae1 WT. ↑ indicating significant upregulation; ↓ indicating significant downregulation; ↔ indicating no difference compared to C57/Bl6 Control, or Ae1 WT respectively.

### ***10.3 Pendrin expression pattern is normal despite incomplete dRTA***

Renal Ae1 deficiency in mice and humans leads to distal renal tubular acidosis (dRTA). The previous results proofed that induction of chronic metabolic acid loads by oral application of  $\text{NH}_4\text{Cl}$  is not sufficient to cause the abnormal expression of Pds observed in the inner medulla of Ae1 KO mice. However, possible negative effects on Pds expression in the inner medulla induced by Ae1 related dRTA cannot be excluded by this experimental model. Therefore, the expression pattern of Pds was investigated in mice deficient for the B1 subunit of V-ATPase (B1) or Rhcg. Both mouse models were reported to develop dRTA upon induced acid-base disturbances (chapter 7.1).

In contrast to Ae1 deficient mice, representative photomicrographs of kidneys from B1 and Rhcg deficient mice did not show signs of abnormal Pds expression in cortex or medulla (Fig. 10.37). Hence, Pds expression pattern in the renal inner medulla is normal despite incomplete dRTA. Moreover, Ae1 and the  $\alpha 4$  and B1 subunits of V-ATPase were not differentially expressed in kidneys of B1 or Rhcg mutant mice compared to WT (data not shown). However, both mouse models were investigated under basal conditions, where acid-base homeostasis seems compensated despite their renal defects.



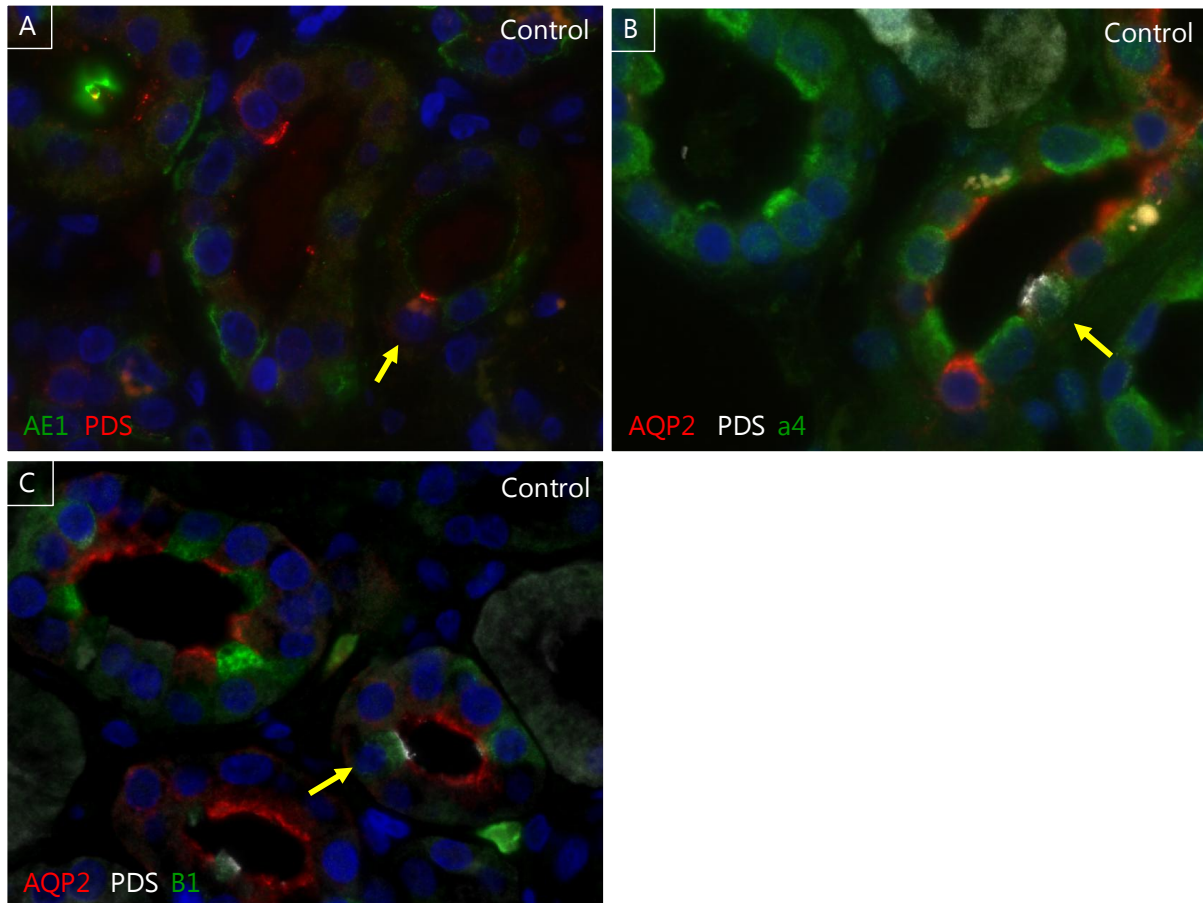
**Fig. 10.37 Immunologic localization of Pds in adult (P84)  $B1^{-/-}$  and  $Rhcg^{-/-}$  mouse kidney.** Representative merged photomicrographs from renal cortex (CCD) of  $B1^{-/-}$  (A) and  $Rhcg^{-/-}$  (C) mice. Representative merged photomicrographs from renal inner medulla (IMCD) of  $B1^{-/-}$  (B) and  $Rhcg^{-/-}$  (D) mice. Individual Aqp2 (aquaporin 2) expression in renal CCD and IMCD identifies principal cells. A, C In the CCD, Pds (pendrin) localizes to the luminal membrane of Aqp2 negative cells, identifying B-ICs. B, D Pds is absent from the mature inner medulla of  $B1^{-/-}$  and  $Rhcg^{-/-}$  mice (n = 3 per genotype). Nucleic acid is stained with DAPI. § indicating Pds positive cell; # indicating Aqp2 positive cell; \* indicating Aqp2/Pds negative cell. Magnification 400x.

#### ***10.4 Expression of acid-base transporters in human Control kidney and a case of AE1 related dRTA***

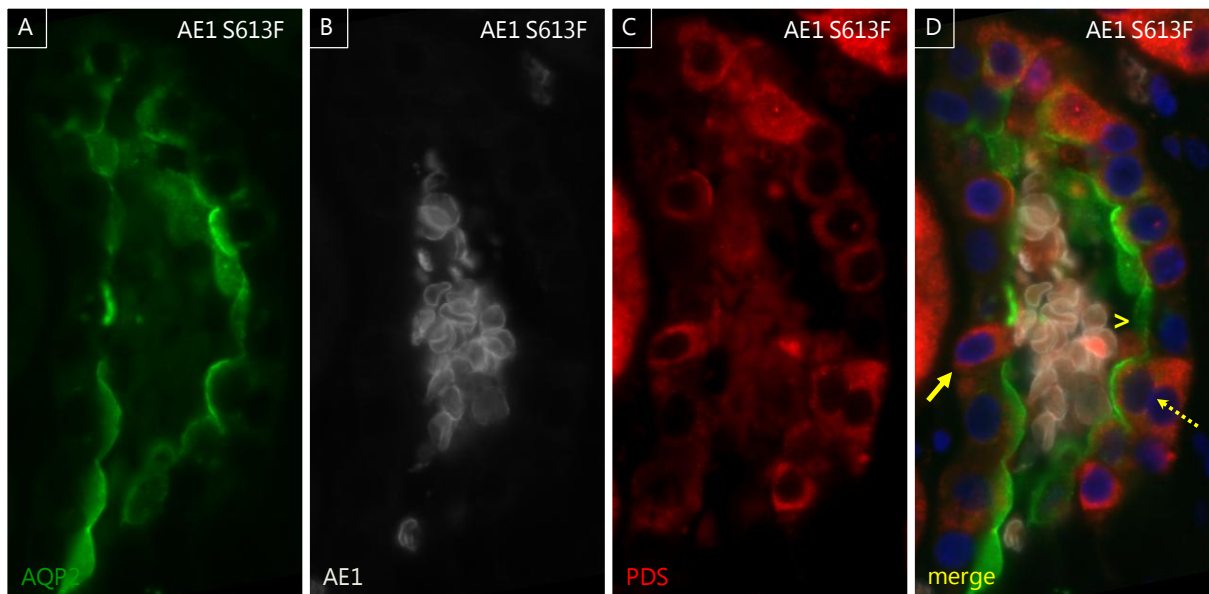
In humans, several mutations of AE1 account for the development of primary dRTA (chapter 4.5.1, Tab. 4.2, chapter 6.1.2). However, these patients are rarely biopsied since diagnosis of disease is made from the clinical presentation and genetic analysis. A renal biopsy of a patient with the dominant negative AE1 S613F mutation (kind gift of Dr. R. Unwin, University College of London, UK) was investigated to search for signs of CD cell abnormalities using immunofluorescent markers. As discussed already in detail, under physiologic conditions luminal expression of PDS labels type B-ICs, whereas basolateral presence of AE1 marks type A-ICs (Fig. 10.38A). Similar to mouse and rat kidneys, AQP2 is expressed on the apical cell membrane of principal cells, while the  $\alpha 4$  and B1 subunit of V-ATPase is expressed in intercalated cells (Fig. 10.38B, C). Consistent with observations in mouse kidney, human carbonic anhydrase II (CA II) expression was found in AQP2 positive and negative cells of the CNT in Control kidneys and is therefore not considered as a IC cell-specific marker (data not shown).

In a renal biopsy of the above mentioned patient with the dominant negative AE1 mutation S613F a reduced abundance of ICs with a concomitant increased presence of PCs was obvious. Variants of the IC type were detected using AE1, PDS and V-ATPase markers ( $\alpha 4$  and B1 subunit) and rarely seen. In contrast, PCs labelled with AQP2 seemed to be the major cell type of the collecting ducts. Furthermore, AE1 immunoreactivity was absent in cells of the connecting tubule and collecting duct in the kidney with AE1 S613F mutation. However as expected, AE1 staining was preserved in erythrocytes (Fig. 10.39B). In addition, PDS positive cells were rarely detected in AE1 S613F renal tissue and PDS seemed to be expressed mostly in the cytosol instead of the apical membrane (Fig. 10.39C). Moreover, few cells coexpressed PDS along with AQP2. Notably, the medullary region was not present in this biopsy as usual. Similarly, the majority of  $\alpha 4$  positive cells unusually coexpressed AQP2 (Fig. 10.40). Additionally, B1 immunoreactivity appeared markedly reduced and B1 positive cells were hardly present (Fig. 10.41). However, because of the single case relative

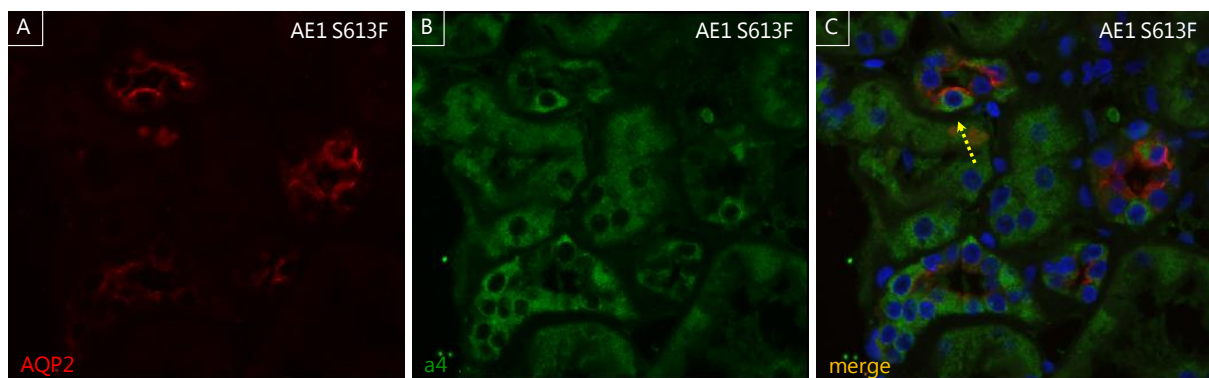
abundance of the labelled cells was not assessed and quantification of relative protein abundance is not possible.



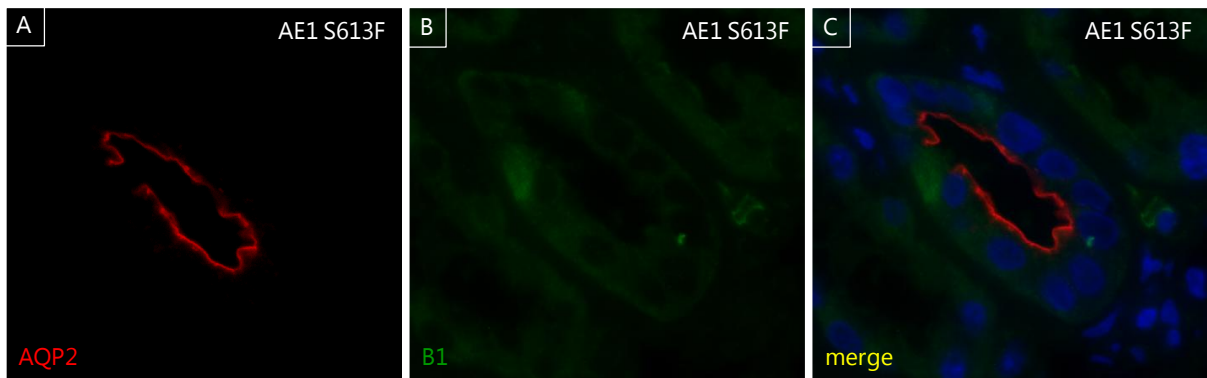
**Fig. 10.38 Expression pattern of collecting duct marker proteins in human Control kidney. A)** AE1 (Anion Exchanger 1) and PDS (Pendrin) are expressed in distinct types of intercalated cells. Basolateral expression of AE1 indicates presence of type A-IC. Apical presence of PDS labels type B-IC (arrow). **B, C)** AQP2 (Aquaporin 2) is exclusively expressed in principal cells. The α4 subunit of V-ATPase is expressed in intercalated cells of the renal collecting duct system and proximal parts of the nephron (**B**). The β1 subunit of V-ATPase is specifically expressed in renal IC (**C**). Colocalization of basolateral V-ATPase with luminal PDS confirms type B-IC identity (arrow) (**B**), while equilateral V-ATPase and PDS expression indicate type non-A/non-B IC phenotype (arrow) (**C**). Magnification 400x.



**Fig. 10.39 Renal expression pattern of PDS and AQP2 in a case of AE1 S613F.** **A)** The majority of cells in the human renal cortex express luminal AQP2 (Aquaporin 2). **B)** Immunoreactivity for AE1 (Anion Exchanger 1) was absent in renal cells of the collecting duct, but is conserved in erythrocytes. **C)** A rather cytosolic expression of PDS (Pendrin) was found in some cells of the renal collecting duct. **D)** merge. Some PDS positive cells remain negative for AQP2 (arrow), while other PDS positive cells seem to coexpress AQP2 (dashed arrow). arrowhead indicates AQP2 positive/PDS negative cell. Magnification 1,000x.



**Fig. 10.40 Renal expression pattern of a4 subunit of V-ATPase and AQP2 in a case of AE1 S613F.** **A)** Apical AQP2 (Aquaporin 2) is present in the majority of renal collecting duct cells. **B)** The a4 subunit of V-ATPase (a4, ATP6V0A4) is expressed in intercalated cells of the renal collecting ducts and in cells of the proximal nephron. **C)** merge. The majority of a4 positive cells are also positive for AQP2 (dashed arrow). Magnification 400x.

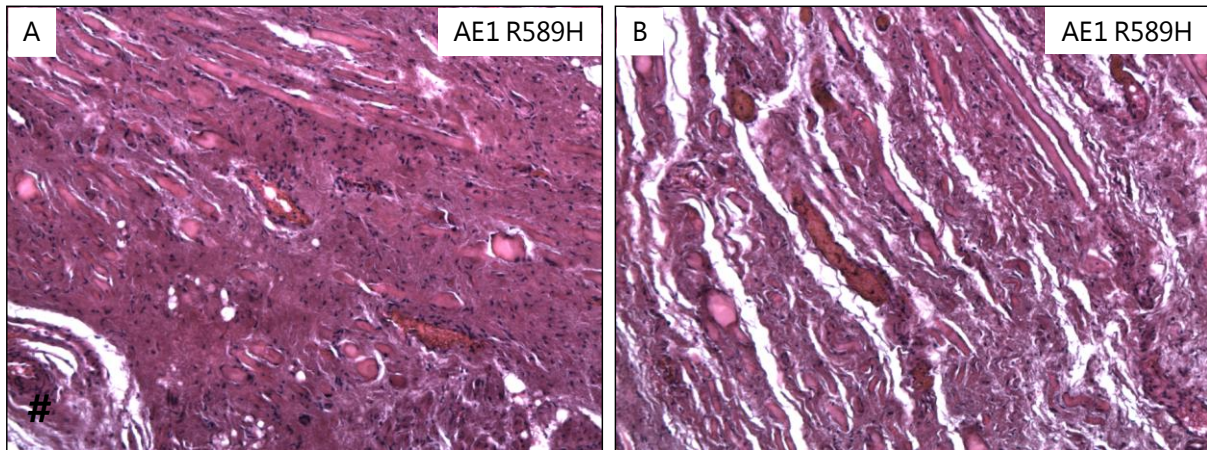


**Fig. 10.41 Renal expression pattern of B1 subunit of V-ATPase and AQP2 in a case of AE1 S613F.**

**A)** Apical AQP2 (Aquaporin 2) is present in the majority of renal collecting duct cells. **B)** The B1 subunit of V-ATPase (B1, ATP6V1B1) is rarely expressed in intercalated cells of the renal collecting ducts. **C)** merge. B1 and AQP2 are expressed in distinct cell types of the renal collecting duct. Magnification 400x.



In another case of primary dRTA related to the dominant negative AE1 R589H mutation (kind gift of Dr. Alper, S.L., Harvard Medical School, Boston, USA) immunofluorescent analysis of the renal CD system failed due to a loss of the renal morphology. Here, chronic nephrocalcinosis and pyelonephritis destroyed the renal tubular structures which were replaced by amorphous fibrotic scars (Fig. 10.42).



**Fig. 10.42 Hematoxylin and Eosin staining of a renal biopsy of a case with AE1 R589H.** The kidney morphology shows amorphous fibrotic remodelling and loss of tubular structures due to nephrocalcinosis and recurring pyelonephritis. # indicates fibrotic remains of a glomerulum. Magnification 100x.

## 11. DISCUSSION

The metanephric mammalian kidney originates from two different primordia, the nephrogenic mesenchymal blastema and the ureteric bud epithelium. The nephrogenic CNT joints the renal CD system to the upstream nephron. The mature renal CD system is the main site of final urinary acidification and according to its functional demands subdivided into the CCD, OMCD and IMCD. However, the renal CNT and CD system are comprised of at least three different cell types, which are unequally distributed (Fig. 2.4). The principal cells (PCs) are the major cell type and involved in systemic electrolyte and water homeostasis. They are interspersed with distinct types of intercalated cells (ICs), which regulate systemic acid-base homeostasis. The two major forms of ICs are the acid secretory type A-ICs and the bicarbonate secretory type B-ICs. In accordance with their function, cells of the renal CD can be distinguished by their ultrastructural appearance and specific marker proteins (Fig. 2.6). In the present study, PCs are identified by the immunologic presence of Aqp2, while both types of ICs are detected by their expression and subcellular localization of the  $\alpha 4$  or B1 subunit of V-ATPase. Furthermore, the two major types of IC are distinguished by their expression of Ae1 (type A-ICs) or Pds (type B-ICs).

The  $\text{Cl}^-/\text{HCO}_3^-$  exchanger Ae1 is essentially expressed in erythrocytes and in the kidney exclusively on the basolateral membrane of type A-ICs. In humans, several mutations of the transporter were identified, which either disturb its function in the erythrocyte or the kidney. Consistent with the role of Ae1 in renal type A-ICs, disturbances of Ae1 expression and function in the kidney lead to distal renal tubular acidosis (dRTA), an inability to acidify the urine. However, data of the present study suggest an additional role for Ae1 in the terminal differentiation of the renal CD system, which is most likely beyond its  $\text{Cl}^-/\text{HCO}_3^-$  exchange.

### ***11.1 Loss of Ae1 coincides with the unusual expression of Pds in the renal inner medulla***

Despite intensive studies of the early events of nephrogenesis and characterization of the adult kidney, only very little is known about the cellular origin of the different cell types of the renal CD system. Studies of fetal mouse, rat and human kidneys suggest the development of PCs and subtypes of ICs from distinct pools of precursor cells in the renal CNT and MCD (chapter 2.5, Fig. 2.5). Interestingly, subtypes of ICs are not detected in the renal CCD before birth. However, with increasing functional demands of the maturing postnatal kidney, the CD system undergoes continuous remodelling resulting in a distinct segmental pattern. In the adult kidney, type B-ICs are exclusively present in the renal CNT and CCD, while the relative abundance of type A-ICs increases towards the inner medulla with a maximum in the outer medullary region (Fig. 2.4, 2.5).

Surprisingly, immunofluorescent investigation of mature kidneys from juvenile (P30) and adult (P84) Ae1 deficient mice revealed an unusual expression of Pds in cells of the inner medulla (Fig. 10.2, 10.6, 10.7). Moreover, some cells coexpressed Pds along with the PC marker Aqp2. Interestingly, the later observation was confined to the inner medulla of mature Ae1 deficient kidneys, as a normal cellular Pds expression was detected in the mature cortex and in embryonic kidneys.

Only the quantification of immunologically counter-labelled cells in juvenile and adult mouse kidneys revealed some differences concerning the cellular distribution among the genotypes. Consistent with chronic dRTA, the relative abundance of Pds<sup>+</sup> cells in the renal CNT and CCD were significantly reduced, while Aqp2<sup>+</sup> cells increased in adult (P84) Ae1 KO kidneys. However, in the renal OMCD of P84 Ae1 KO mice the relative abundance of Aqp2<sup>+</sup>/Pds<sup>+</sup> coexpressing cells increased, probably on the expense of unlabelled cells. In addition, the relative abundance of Pds<sup>+</sup> and Aqp2<sup>+</sup>/Pds<sup>+</sup> coexpressing cells in the renal ICMD of P84 Ae1 KO mice increased with a modest decrease in unlabelled cells.

In contrast, Aqp2 and Pds expressing cells in the CNT of juvenile (P30) mouse kidneys were not altered between WT and Ae1 KO mice. Moreover, Pds<sup>+</sup> cells increased in abundance on the expense of unlabelled cells in the CCD and OMCD of P30 Ae1 KO mice. Likewise, Pds<sup>+</sup> and Aqp2<sup>+</sup>/Pds<sup>+</sup> coexpressing cells in the renal ICMD of P30 Ae1 KO mice increased, while unlabelled cells modestly decreased. Strikingly, the unusual expression of Pds in Ae1 deficient IMCDs is accompanied by a reduced relative abundance of Aqp2<sup>+</sup> in P30 mice, while it is paralleled by a reduced abundance of unlabelled cells in P84 kidneys.

The altered cellular expression pattern between P30 and P84 Ae1 KO mice might be a consequence of two circumstances. The juvenile mice just experienced a dietary switch from mother milk to rodent chow, which was shown to provide a strong net alkali load [227]. Consistently, the urinary pH of juvenile mice after weaning from the mother milk was shown to be more alkaline [31]. Hence, the increased need to excrete excess base equivalents from the body might reflect the unexpected increased presence of Pds<sup>+</sup> cells in the CCD and OMCD of P30 Ae1 KO kidney despite early dRTA. It remains speculative, whether this is due to a delayed apoptotic removal of Pds<sup>+</sup> cells from the MCD or an increased proliferation rate of Pds<sup>+</sup> cells. In addition, adult (P84) Ae1 KO mice suffered from chronic dRTA, which affects renal water handling [190, 218]. This might also be the reason for the reduced number of Aqp2<sup>+</sup> in the renal CNT of juvenile (P30) WT mice compared to adult (P84) WT animals. Furthermore, an adaptive increase of Aqp2<sup>+</sup> cells upon prolonged dRTA might mask the abnormal differentiation of Aqp2<sup>+</sup> cells in the renal IMCD, or drive the abnormal differentiation of cells other than Aqp2<sup>+</sup>. However, in kidneys of juvenile Ae1 KO mice, which might have a less pronounced dRTA, the unusual expression of Pds in the IMCD seems to origin from the abnormal differentiation of cells retaining or acquiring Aqp2 expression.

Consistent with the observations in Ae1 deficient mouse kidney, PDS expression seemed altered in a renal biopsy of a patient with primary dRTA due to the dominant negative AE1 S613F mutation (chapter 10.4). This mutation is known to exclusively affect the expression and functionality of AE1 in renal type A-ICs, but not in erythrocytes. *In vitro* experiments using polarized MDCK cells suggest that the AE1 S613F mutant protein is retained in the ER and thus does not reach the basolateral cell membrane of type A-ICs. Since reticulocytes are not polarized and have different chaperones and trafficking mechanisms than renal epithelial cells, they might circumvent the dominant negative effect of the mutant and escape early ER degradation.

However, a previous investigation of the patient's renal biopsy documented a reduced number of renal ICs and a solely cytosolic expression of AE1 S613F, with a preserved expression in erythrocytes [138]. Similar results were obtained using immunofluorescent molecular CD cell markers in the present study. The abundance of renal ICs (identified by V-ATPase, PDS or AE1) was remarkably low and PCs (identified by AQP2) seemed to comprise the major cell type of the renal CD system. In the present study, AE1 immunoreactivity was only detected in erythrocytes but the accessibility of the renal epitope might have been lost over the years of storage, especially since the original work mentioned a rather weak AE1 signal intensity in renal epithelial cells [138]. Anyway, V-ATPase and PDS expression were detected only in very few cells of the renal CD system and PDS localization appeared to be mostly cytosolic. Notably, the CD section represented on the biopsy were exclusively from the CNT and CCD segment as assessed by the gross morphology of the surrounding tissue and given by the usual technique to take biopsies.

Similarly, a reduced number of renal ICs from the CD system was reported for a case with the dominant negative AE1 R589H mutation [208].

Hence, Ae1 deficiency in mouse and man seems to correlate with abnormal subcellular expression of pendrin and severely reduced number of ICs.

### ***11.2 The type A-IC phenotype is preserved in Ae1 deficient kidneys***

In order to further elucidate which CD cell type might be abnormally differentiated in Ae1 KO kidneys, the expression pattern of V-ATPase was examined in P30 mice. The expression pattern of the kidney specific  $\alpha 4$  subunit of V-ATPase did not show any significant differences between the genotypes. However, the IC-specific B1 subunit of V-ATPase partially reflected the unusual presence of Pds in the renal MCDs of Ae1 deficient mice. Here, an increased abundance of  $B1^+/Aqp2^+$  coexpressing cells was observed in Ae1 KO kidneys, which reached significant difference in the IMCD of P30 mice. In line with the previous observation,  $Aqp2^+$  cells of the Ae1 deficient IMCD were simultaneously decreased. Therefore, some  $Aqp2$  expressing cells of the IMCD of Ae1 KO mice might either coexpress Pds ( $Aqp2^+/Pds^+$ ) or B1 ( $Aqp2^+/B1^+$ ) or both ( $Aqp2^+/Pds^+/B1^+$ ), respectively. The unchanged expression pattern of individually  $B1^+$  cells in Ae1 deficient kidneys suggests a normal differentiation of the type A-IC phenotype, which certainly represents the major IC subtype in the pool of unlabelled cells. This observation implies that renal Ae1 deficiency does not necessarily result in a loss of type A-ICs, but rather a loss of a proper molecular marker. Unfortunately, no other unequivocal marker of type A-IC is available to date to further confirm maintenance of this cell type. However, the unusual expression pattern of Pds expressing cells in Ae1 KO kidneys might origin from the abnormal differentiation of PCs and affects the type B-IC phenotype or the type non-A/non-B IC variant. However, further experiments are needed to answer this question.

### **11.3 *Foxi1* expression is altered in *Ae1* deficient kidneys**

Transcription factors of the forkhead box family are involved in early nephrogenesis and terminal differentiation of the renal CD system (chapter 2.7.3). *Foxi1* deficient mice were reported to lack molecular markers of renal ICs and consequently develop dRTA [83]. Moreover, the renal CD system of *Foxi1* KO mice consists of a single cell type which coexpresses *Aqp2* and *CaII* (Fig. 2.11). Similarly, *Foxi1* homologues are required for epidermal ionocyte differentiation in larvae of *Xenopus laevis* and zebrafish (Fig. 2.7, 2.8) [51, 54, 56]. Therefore, the *Foxi1* expression pattern was examined in *Ae1* deficient kidneys by immunohistochemistry. Previous studies mentioned *Foxi1* expression in the murine distal tubule and more precisely in type A-ICs and B-ICs, but not in PCs [82, 83]. However, the preliminary results of the present study reveal a nuclear expression of *Foxi1* mainly in  $B1^+$  ICs, but also in few  $Aqp2^+$  PCs (Fig. 10.11-16). The contrary cellular localization of *Foxi1* with previous reports might be due to the distinct experimental approach. Anyhow, quantification of the *Foxi1* expression pattern in *Ae1* WT and KO kidneys of juvenile mice revealed some interesting differences. The relative abundance of individual  $Foxi1^+$  cells was significantly increased in the renal IMCDs of *Ae1* KO mice compared to WT. Moreover, a similar amount of  $Aqp2^+/Foxi1^+$  cells was increased in IMCDs of *Ae1* KO mice, even though it did not reach statistical significance. Furthermore,  $Aqp2^+/Foxi1^+$  cells in the CCD of *Ae1* deficient kidneys were significantly increased, while the relative abundance of  $Foxi1^+$  and unlabelled cells was simultaneously reduced. Similarly,  $Pds^+$  cells were increased in the CCD of *Ae1* deficient kidneys on the expense of unlabelled cells, while the relative abundance of  $Aqp2^+/B1^+$  tends to increase on the expense of  $Aqp2^+$  cells. Likewise, the increase of  $Pds^+$  and  $Aqp2^+/Pds^+$  cells in the renal IMCD of *Ae1* KO mice on the expense of  $Aqp2^+$  cells is paralleled by an increase in  $Foxi1^+$  cells. Although the percentage of  $Aqp2^+/Foxi1^+$  cells is smaller than the number of cells with unusual *Pds* and *B1* expression, their concomitant appearance is likely related to each other and thus caused by *Ae1*

deficiency. Moreover, *Foxi1* is a known positive transcriptional regulator of the B1 subunit of V-ATPase, *Ae1* and *Pds* [83, 85, 87].

Interestingly, mice with a conditionally disrupted Notch signalling pathway in the renal CD system suffer from nephrogenic diabetes insipidus based on the loss of *Aqp2* expressing PCs (chapter 2.7.2) [77]. Additionally, an increased abundance of V-ATPase expressing cells with a concomitant increased mRNA expression of *V-ATPase*, *Ae1*, *Pds* and *Foxi1* was noted. Consistently, a mouse model with a constitutive activation of the Notch signalling pathway in the renal CD system was devoid of molecular markers for ICs and expressed only *Aqp2* positive PCs (Fig. 2.10). Hence, terminal differentiation of the renal CD cells is regulated by the Notch signalling pathway and *Foxi1*, but the definite interaction of these transcriptional regulators is not yet elucidated. Studying the expression of the molecular players of the Notch signalling pathway during collecting duct development in WT mice and its regulation in *Ae1* KO kidneys would add further important information.



#### **11.4 *Cp2l1* expression is altered in *Ae1* deficient kidneys**

Similar to Foxi1 KO mice and those with disrupted Notch signalling pathway, animals deficient for the transcription factor Cp2l1 are devoid of IC marker in the renal CD system (chapter 2.7.4, Fig. 2.11) [90]. The role of Cp2l1 in the mammalian kidney is only poorly investigated, but studies of the epidermal development of ionocytes in *Xenopus* and zebrafish larvae suggest a regulatory role of Cp2l1 homologues in a Pds expressing ionocyte subtype [51, 57]. Moreover, the transcriptional control of Cp2l1 involves the interaction with the Notch signalling pathway and Foxi transcription factors (Fig. 2.7, 2.8). However, whether this is also true for Cp2l1 in murine kidney cells is not known. Preliminary data from immunologic localization studies on P30 WT mouse kidneys show coexpression of nuclear Cp2l1 mainly with Aqp2 positive and less with Aqp2 negative cells along the renal CD system (Fig. 10.17, 10.18). Apparently, the abundance of Cp2l1 positive cells increases from the CNT towards the inner medulla, where Cp2l1 seems to be expressed in almost every cell. Similarly, the presence of Aqp2 expressing PCs increases from the CNT towards the inner medulla. In addition and in line with a previous report, nuclear Cp2l1 expression is detected in some tubuli of the DCT and TAL [90]. Even though detailed colocalization studies are not yet available, a striking difference in the expression of Cp2l1 was observed in *Ae1* deficient kidneys compared to wildtype. The nuclear expression profile of Cp2l1 in inner medullary collecting duct cells was completely absent and moreover appeared to be shifted to the apical membrane in some cells. Whether the altered subcellular localization is related to different activity states of the transcription factor needs to be investigated. However, the loss of Cp2l1 from the nuclei of IMCD cells seems to coincide with the abnormal cell differentiation described upon renal *Ae1* deficiency.

### **11.5 Late onset of *Ae1* related abnormal differentiation of renal IMCD**

Although Pds expressing cells in kidneys of *Ae1* deficient embryos seemed to be preserved, an abnormal regulation of transporters and transcription factors involved in CD cell function and specificity cannot be ruled out. Therefore, mRNA expression levels were compared at different developmental ages in kidneys of *Ae1* mutant mice (Tab. 10.20). Interestingly, alterations of genes implicated in renal acid-base handling (*Ae1*, *Pds*, *B1 subunit of V-ATPase* and *Rhcg*) were not altered before postnatal day 30 (P30). This result supports the afore mentioned impact of a dietary switch from mother milk to rodent chow on systemic acid-base homeostasis and the concomitant mutual reaction with *Ae1* related dRTA. Alternatively, it may reflect the completion of postnatal maturation of the kidney which includes the removal of Pds<sup>+</sup> cells from the medulla. Surprisingly, mRNA abundance of *Foxi1* and *Cp2l1* were not different between *Ae1* WT and mutant mice at any time point of life investigated, despite the marked changes in the protein expression pattern. However, gene expression does not necessarily correlate with protein abundance or activity. On the other hand, genes suggested to be involved in the p53 signalling cascade and probably developmental fate of renal CD cells (*Gdf15*, *Klf4* and *Bdkrb2*) were steadily upregulated in *Ae1* KO mice after postnatal day 3 (P3).

Together with the immunofluorescent analysis of molecular IC markers, the data suggest a late onset of *Ae1* related abnormal terminal differentiation of renal CD cells which might be mutually affected by systemic or renal acid-base dysregulation. However, the impact of dRTA seems to be confined to the renal CCD of adult (P84) mouse kidneys, where Pds expressing cells were reduced and Aqp2 positive PCs increased. In addition, the loss of cellular *Ae1* expression seems to induce the abnormal differentiation of Aqp2<sup>+</sup>/Pds<sup>+</sup> coexpressing cells, particularly in the renal IMCD. Similarly, a late onset of IC depletion was reported in *CaII* deficient mice (chapter 2.7.6, Fig. 2.15) [103].

### ***11.6 Chronic metabolic acidosis induces adaptive remodelling of the distal renal tubule but does not affect terminal differentiation of renal CD cells***

Disturbances in systemic acid-base homeostasis are known to induce adaptive responses of the kidney, including a remodelling of the renal tubule (chapter 5). Furthermore, some inherited or acquired defects of renal acid-base transport go along with renal tubular acidosis. Ae1 deficient individuals develop chronic dRTA and accordingly Pds expressing type B-ICs were markedly reduced in adult (P84) Ae1 KO mice. Similarly, ICs in the renal CNT and CCD of patients with AE1 related primary dRTA appeared smaller in size and reduced in their abundance. In order to investigate the impact of chronic acid-base disturbance on the terminal differentiation of the renal CD system, immunofluorescent analysis and mRNA expression levels were performed on kidneys of mice with induced chronic metabolic acidosis for 8 weeks. As expected and consistent with the observations in P84 Ae1 deficient mice, kidneys of chronically acid loaded mice showed a reduced abundance of Pds expressing type B-ICs in the CCD, while Aqp2<sup>+</sup> cells increased. However and in contrast to P84 Ae1 KO mice, the relative abundance of Aqp2<sup>+</sup> cells decreased in the renal OMCD of acidotic animals, while unlabelled cells increased concomitantly. Likewise in contrast to Ae1 deficient kidneys, CD cells coexpressing markers of ICs and PCs respectively were not detected in NH<sub>4</sub>Cl treated mouse kidneys. Co-immunolabelling using further molecular markers for distinct types of ICs (V-ATPase subunits and Ae1), confirmed the increase of Aqp2 expressing PCs on the expense of Pds<sup>+</sup> type B-ICs in the renal CCD of acidotic mice (Fig. 10.25 to 10.28). In addition, assessment of the relative protein abundance of Aqp2, Pds, Ae1 and B1 subunit of V-ATPase by western blotting revealed comparable results (Fig. 10.29 to 10.32). The relative protein abundance of Aqp2 was increased in NH<sub>4</sub>Cl treated kidneys compared to Control, while Pds abundance was significantly reduced. Furthermore and expected, the relative abundance of Ae1 and the B1 subunit of V-ATPase was not altered upon chronic acid loading.

Finally, the renal mRNA expression pattern of acid-base transporters and transcriptional regulators was compared between  $\text{NH}_4\text{Cl}$  loaded mice and *Ae1* deficient mice (Fig. 10.35, 10.36). As illustrated in the simplified table of Fig. 10.36, chronic metabolic acidosis induces a completely distinct adaptive regulation of acid-base transporters and transcriptional regulators compared to *Ae1* related dRTA. Consistent with previous observations,  $\text{NH}_4\text{Cl}$  treatment induces an upregulation of *Ae1* mRNA levels with a concomitant downregulation of *Pds* mRNA. In contrast, *Pds* mRNA expression levels are not different in kidneys of *Ae1* WT and KO mice. Interestingly and likewise in contrast to *Ae1* KO mice, renal *Cp2l1* mRNA expression was significantly upregulated in  $\text{NH}_4\text{Cl}$  treated mice compared to Control. Moreover, *Gdf15* and *Klf4* mRNA expression was significantly upregulated in *Ae1* deficient kidneys compared to *Ae1* WT, while the transcription of target genes of the p53 signalling pathway were not different in Control and  $\text{NH}_4\text{Cl}$  loaded mice.

Furthermore, western blotting of Gdf15 showed significant downregulation of a probably premature variant, while the supposedly secreted and molecularly active form of Gdf15 is significantly upregulated in  $\text{NH}_4\text{Cl}$  treated mouse kidney (Fig. 10.33). In contrast, Gdf15 related variants appeared not regulated in *Ae1* deficient mice. Whether the secreted active signalling peptide of Gdf15 is also represented and regulated in the blood of the mice remains speculative. Gdf15 is a multivariant molecule and involved in several distinct cellular responses. Gdf15 has been reported, together with p53 and PI3/Akt/mTOR signalling cascade, to be involved in the early proliferation of particularly type A-ICs upon induced metabolic acidosis [146]. The present study is the first one that shows the protein pattern of renal Gdf15 detected on western blot, since previous studies were based on mRNA expression. More strikingly, our study suggests for the first time a role for Gdf15 in renal adaptation during chronic acid-base disturbance. Cell proliferation events are rather unlikely in the compensated late phase of chronic acidosis. Hence, Gdf15 might rather play a role in reducing the cellular stress induced by prolonged acidity and may promote anti-apoptotic actions. Such anti-apoptotic actions might be necessary to maintain

the unusual high abundance of Aqp2 expressing PCs in the CCD of acidotic kidneys. Similarly, pro-apoptotic actions of Gdf15 could be necessary to mediate or maintain the reduced presence of Pds expressing type B-ICs in acidotic renal CCDs. However, further experiments investigating the nature of Gdf15 and the related molecular variants detected in the kidney are necessary to determine its precise role in acid-base disturbances.

*Foxi1* mRNA expression levels were not altered in chronic metabolic acidosis or dRTA. Anyhow, quantification of labelled cells by immunofluorescence revealed differences in the cellular distribution of Foxi1 in Ae1 KO kidneys compared to WT. Detection of Foxi1 by western blot on kidney samples derived from NH<sub>4</sub>Cl loaded and Control mice, demonstrated the presence of distinct Foxi1 related bands which differed markedly in size (Fig. 10.34). Even though the Foxi1 related bands did not appear to be regulated upon acid-base disturbance, their discovery is meaningful not at least because they are detected for the first time. In nuclear protein extracts, three Foxi1 related bands were detected, of which only one corresponds to the expected molecular weight. In contrast, the expected Foxi1 band was absent in cytosolic fractions of the respective kidney samples, while the two Foxi1 related bands of higher molecular weight were present. This finding may suggest, that the Foxi1 variant with the expected molecular weight is the transcriptionally active monomer and therefore present in the nuclear protein fraction but not in the cytosol. Likewise the two Foxi1 related bands of higher molecular weight might be distinct homomultimers or heteromultimers which translocate from the cytosol into the nucleus upon activation or inactivation. Again, further experiments are necessary to proof this idea and characterize the signalling network of Foxi1 in renal CD cells in detail. In addition, immunofluorescent analysis and quantification of the Foxi1 expression pattern for kidneys of Control and NH<sub>4</sub>Cl treated mice are missing.

However, altogether the data presented in this study suggest, that prolonged metabolic acidosis induces alterations of the cellular composition of the renal CD system which favour the renal excretion of excess acid along with increased water retention, but does not alter the distinction between differentiated CD cell types.

### ***11.7 Terminal differentiation of renal CD cells is not affected by incomplete dRTA***

Inherited dRTA is characterized by the inability to acidify the urine and hence a primary renal defect. In contrast, induced metabolic acidosis in mouse models is a forced disturbance mediated by artificial oral acid loads. Therefore, the two acid-base disturbances might somehow result in different renal adaptive responses and are difficult to compare. Thus, two more distinct transgenic mouse models were examined for the impact of renal acid-base disturbance on CD cell differentiation. Mice deficient for the B1 subunit of V-ATPase, as well as mice lacking Rhcg were shown to develop dRTA when challenged with increased acid loads [114, 215]. Because these mice do not show symptoms of dRTA under basal conditions, adaptive mechanisms such as altered CD cell composition, might be activated and compensate the obvious renal defect.

Despite incomplete dRTA and in contrast to Ae1 deficiency, both mouse models did not show any signs of abnormal expression of Pds, Ae1, a4 and B1 subunits of V-ATPase when investigated by immunofluorescence (Fig. 10.37).

Likewise, Pds deficient mice were not reported to suffer from abnormal CD cell differentiation, despite their severe renal defect [213].

Therefore, our data suggest that the loss of Ae1 protein in the kidney affects the terminal differentiation of cells in the renal IMCD, inducing an abnormal expression of Pds and moreover a coexpression of Pds and Aqp2 independently of the acid-base disturbance.

### **11.8 Hensin is not expressed in mouse renal tissue**

Over the past two decades, the extracellular matrix protein hensin was continuously suggested to play a crucial role in the terminal differentiation of renal CD cells and recently a hensin deficient mouse model was reported to develop dRTA [99, 100]. Accordingly, we set out to investigate a possible dysregulation of hensin in Ae1 deficient mouse kidney. However, in the present study we failed to detect mRNA expression of *hensin* in premature and mature kidneys of mice (Fig. 10.21).

Hensin was originally discovered *in vitro*, in immortalized cells derived from IMCDs of rabbit kidney. Furthermore, *in vitro* experiments using this cell model suggested a role for hensin in the transdifferentiation of type B-ICs to type A-ICs and *vice versa*. In this experimental protocol, a mixed population of rabbit IMCD cells was cultured with different initial density, which seemed to affect and alter the identity of cells (A-IC or B-IC). Hensin was discovered and reported to mediate the molecular switch that determines the differentiation of cells and thus their identity as type A-ICs or B-ICs. However, in a similar cell line derived from renal mouse IMCDs neither *hensin* mRNA was detected nor mRNA of *Ae1*, *Pds* and *V-ATPase* (data not shown), suggesting that the cell population did not contain any cells resembling the phenotype of type A-ICs or type B-ICs. Anyhow, we repeated the experimental protocol as denoted in the original publications, but were still unable to detect any mRNA for molecular IC markers and *hensin* in mouse IMCD3 cells (data not shown). The repetitive negative result is consistent with the known technical limitation of cultured renal CD derived cells, which immediately lose the phenotypic characteristics of ICs and dedifferentiate *in vitro*. However, species differences might explain the opposing results from *in vitro* studies, since the renal CD system of rabbits is slightly different compared to that of mouse, rat or human.

Notably and in spite of the reported phenotype of the renal CD specific hensin KO mouse and its supposedly crucial role in renal IC differentiation, hensin expression has never been shown in renal tissue of mice. In addition, the data presenting the phenotype of the hensin KO mice are not convincing and raise reasonable doubts on the existence and impact of hensin, at least in mouse kidney.

### ***11.8 Ae1 expression is required to maintain terminal differentiation of renal CD cells***

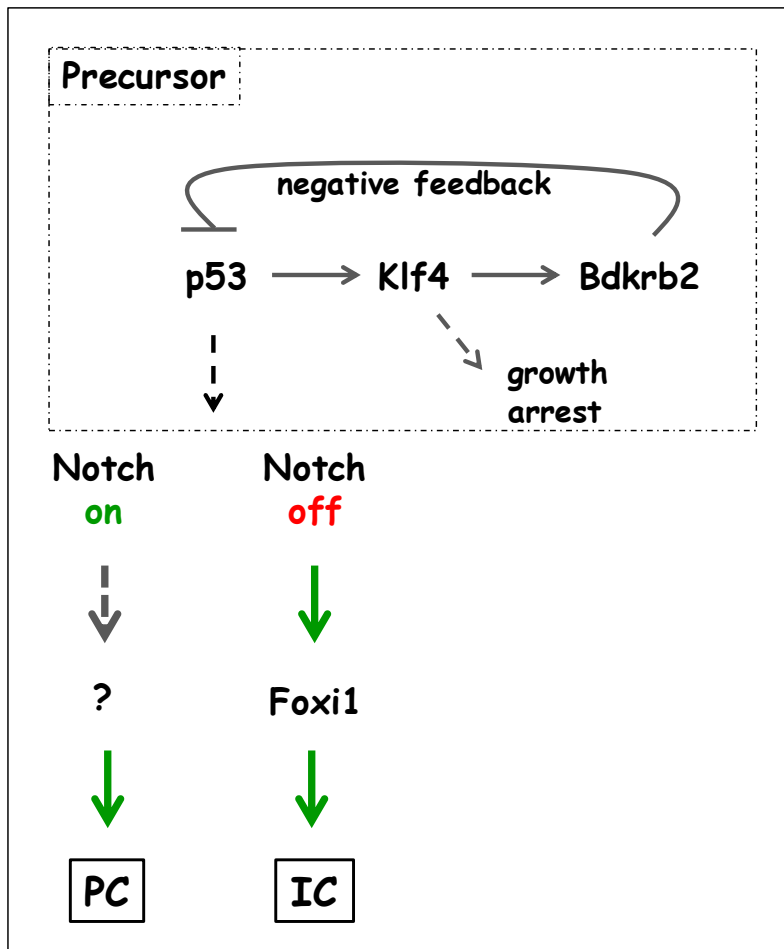
The origin of individual cell types of the renal collecting ducts is not yet entirely proven. But some studies suggest that principal cells and types of intercalated cells origin from the same precursor population, simultaneously in the renal CNT and the inner medullary region [18, 25, 27]. In addition, the transcriptional network and other regulatory signalling molecules involved in terminal differentiation of the renal CDs are only poorly understood. Although some molecules were identified in distinct model organisms, the interaction of the distinct signalling pathways and their cellular localization in mouse kidney is still unknown. Therefore, it is very difficult to get a clear picture of the necessary events and key molecules of renal terminal differentiation and to put all the different information together in the right context. Especially, since it needs to account for the complexity of the cellular events in distinct renal cell populations, the putative interaction of different cell types and signalling networks, and the different systemic and cellular functional demands the maturing kidney is requested to fulfil.



However, the present study provides some new findings regarding a novel and unusual role of Ae1 in the terminal differentiation of renal CD cells, which is most likely beyond  $\text{Cl}^-/\text{HCO}_3^-$  exchange. Ae1 expression in the murine kidney might be a key molecule that triggers the maintenance of terminal differentiation of CD cells. Our findings offer interesting possibilities to fill some of the open gaps of our knowledge and to put important pieces of the puzzle into place. Moreover, we show for the first time the cellular localization of the transcription factors Foxi1 and Cp2l1 in the murine kidney. Furthermore, we provide data indicating a local dysregulation of Foxi1 and Cp2l1 upon renal Ae1 deficiency.

Based on the work by other research groups the following hypothetical model is suggested, summarizing the possible role of some key regulatory molecules in the terminal differentiation of renal CD cells (Fig. 11.1). The cell cycle and transcriptional regulator p53 is expressed in the differentiating zone of the late embryonic kidney and was suggested to influence cell lineage differentiation (chapter 2.7.1, Fig. 2.9). p53 was found to interact with other transcriptional regulators, like Klf4 and Bdkrb2 and to regulate the transcription of renal function genes, such as Aqp2 and  $\text{Na}^+/\text{K}^+$ -ATPase [65]. Recently, the Notch signalling pathway was elegantly shown to determine the cell lineage of renal PC and ICs in the mouse [77]. It is not directly proven for the kidney, but in other tissues p53 family members can modulate the activity of Notch, suggesting a crosstalk of both pathways during terminal CD differentiation [74]. Constitutive activation of the Notch signalling cascade drives the expression of PCs in the absence of ICs. Whether another key regulator is intermediately necessary is not known. On the other hand, constitutive inactivation of Notch leads to the loss of PCs and favours the expression of molecular IC markers, such as V-ATPase, Ae1 and Pds. Furthermore, *Foxi1* mRNA expression levels were significantly increased, suggesting a regulatory interaction of the Notch signalling cascade with Foxi1. Accordingly, Foxi1 mediates the transcriptional activation of molecular IC markers. These observations were also made in epidermal ionocyte

differentiation of *Xenopus laevis* and zebrafish larvae suggesting a well conserved signalling pathway (Fig. 2.7, 2.8) [51, 54].



**Fig. 11.1 Proposed model of CD cell lineage decision in mouse kidney.** Principal cells (PCs) and intercalated cells (ICs) may develop from a common renal precursor. As suggested by El-Dahr, S. and colleagues [66], the cell cycle and transcriptional regulator p53 initiates developmental cell differentiation and requires the interaction with Klf4 and Bdkrb2 (Fig. 2.9). Furthermore, p53 may regulate the Notch signalling pathway promoting cell lineage decisions. The activity of Notch determines the later CD cell type. Constitutive activation of Notch favours the differentiation of PCs in the absence of ICs, while inactivated Notch leads to the differentiation of ICs in the absence of PCs [77]. Whether another transcriptional regulator is involved in the differentiation of PCs is not yet known. However, Foxi1 is essentially involved in the terminal differentiation of the IC phenotype [83].

In the murine kidney, loss of the transcription factors Foxi1 and Cp2l1 goes along with the loss of *Ae1*, *Pds* and *V-ATPase* [83, 90]. Moreover, Foxi1 was directly shown to be a transcriptional regulator of *V-ATPase*, *Pds* and *Ae1* [85, 86]. However, whether Foxi1 and Cp2l1 are only expressed in the developing kidney or also during adulthood had not been known and shown in the present study. Moreover, the CD cell type expressing Foxi1 and Cp2l1 was uncertain and has been analyzed to some extent in this work. The present study shows for the first time, that Foxi1 and Cp2l1 are expressed in distinct CD cell populations of the mature mouse kidney. As suggested previously, Foxi1 expression was mainly found in V-ATPase expressing ICs along the renal CD system. This is consistent with its role as a transcriptional regulator of specific IC transporters. However, we found that Cp2l1 expression colocalized mostly with Aqp2 expressing principal cells. This is surprising, since Cp2l1 deficiency also correlates with the loss of molecular IC markers, rather suggesting its expression in the same locus than its target genes. However, Cp2l1 is an unusual member of the grainyhead family, because it acts as a transcriptional repressor. Interestingly, observations in the epidermis of amphibians suggest an interaction of Notch, Foxi1 and Cp2l1 (Fig. 2.7, 2.8). Hence, Cp2l1 in the mouse kidney might be the proposed intermediate regulator in the cell lineage differentiation of PCs, similar to Foxi1 in the IC lineage (Fig. 11.1). Accordingly, Cp2l1 might modulate the activity of Notch in adjacent cells, which results in the activation of Foxi1 and thus differentiation of IC phenotype. The expression pattern of both transcription factors, Foxi1 and Cp2l1, was altered in *Ae1* deficient mature kidneys as already discussed. These observations suggest that Foxi1, Cp2l1 and *Ae1* somehow interact together in renal CD cells, particularly in the inner medulla. A negative regulation by some unknown molecule or a negative autoregulation of Cp2l1 might be necessary in murine kidney, similarly to observations made in zebrafish epidermal development (Fig. 2.8). Accordingly, the loss of nucleic Cp2l1 in some cells of *Ae1* deficient renal IMCDs might prohibit a repression of Aqp2 or might prevent a negative regulation of Foxi1 which promotes the expression of *Pds* in these cells. Consequently, Foxi1 positive cells were more abundant in *Ae1* deficient renal IMCDs (Fig. 10.13D, 10.16D)

leading to the unusual coexpression of Aqp2 and Pds. However, a definite proof is pending and further investigations are necessary.

With respect to our investigations of incomplete dRTA mouse models, the induced chronic metabolic acidosis model and the fact that Pds KO mice do not have abnormally differentiated CD cells, we can conclude that the presence of Ae1 protein is required in the terminal cell differentiation. Respectively, reports by others and our immunofluorescence data from patients with dRTA related AE1 mutations suggest that the proper expression and subcellular localization of the protein is primarily required and not its anion exchanger activity. Moreover, the very preliminary human data suggest that the findings from the mouse model may be transferable to the human situation.

Ae1 is known to form multi-protein complexes in the erythrocyte, which is essential for the cytoskeletal integrity and transport function (chapter 6.1.2) [186]. Similarly, the kidney specific variant of Ae1 is suggested to interact directly or indirectly with distinct proteins, such as ILK (integrin-linked kinase) and Glyceraldehyde 3-phosphate dehydrogenase (GAPDH) [228-230]. Moreover, Ae1 was shown to form a transport metabolon together with CaII in erythrocytes and the kidney enhancing the transport activity [197]. Likewise, the Ae1-CaII transport metabolon might generate a local cellular pH microenvironment that influences the activity of vicinal regulatory proteins thereby triggering signalling pathways [231].

Deficiency of CaII results in the depletion of ICs from the renal CD system with a late onset in the mature mouse kidney (chapter 2.7.6) [101, 103]. Consistently, pharmacologic inhibition of CaII with acetazolamide reduces the abundance of ICs and alters their morphology [104]. Interestingly, the CD system of Foxi1 deficient mice did not only lack ICs, but consisted of a single cell type which coexpressed Aqp2 and CaII (Fig. 2.11) [83]. Indeed we also report here that CaII expression appears normal in Aqp2 expressing cells of the CNT and early CCD of mouse and human kidney (data not shown). But we did not find CaII and Aqp2 coexpressing cells in the OMCD and

IMCD. However, the observation in Foxi1 KO mice implicates that the restrictive expression of CaII to ICs of the MCD segments is somehow influenced by Foxi1, even though CAII is not a direct transcriptional target of Foxi1. Hence, Foxi1 might interact with another regulatory molecule or a signalling pathway to mediate a cross talk between subpopulations of CD cells and indirectly suppress the expression of CaII in Aqp2<sup>+</sup> principal cells.

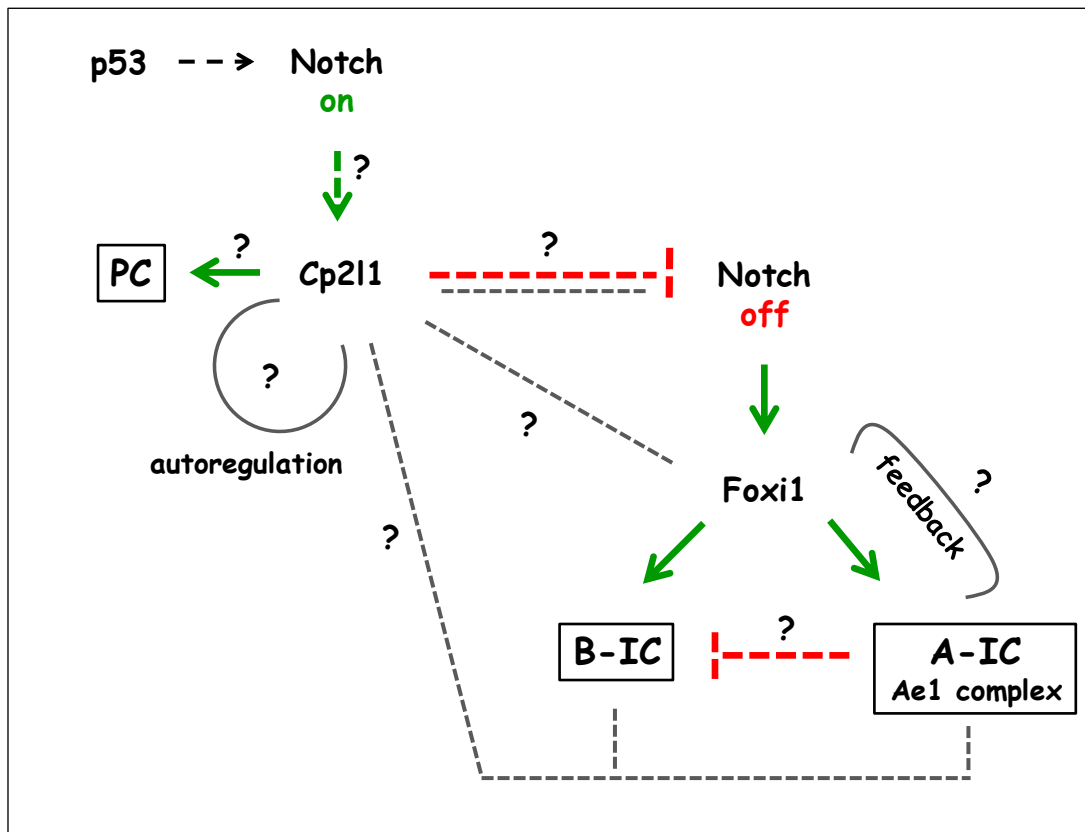
Fox factors were reported to unusually interact with proteins other than transcription factors [232]. Hence, Foxi1 might unusually interact with a multi-protein complex of Ae1 or with Ae1 directly. This speculated interaction could be necessary to maintain or conduct terminal cell differentiation of renal ICs, at least in the mature kidney. A proposed model integrating this speculation and a possible role of the transcriptional repressor Cp2l1 in renal CD differentiation is depicted in Fig. 11.2. In the developing mouse kidney the expression of Pds is observed much later than that of V-ATPase and Ae1 (Fig. 2.5). Hence, type B-ICs develop after type A-ICs. Neither an interconversion of IC phenotypes is proven, nor the question for a common precursor of ICs. However, the presence of a certain type of IC might influence the identity of a neighbouring cell, maybe by paracrine signalling pathways, such as Notch mediated lateral inhibition. This could explain why type B-ICs appear after type A-ICs and our observations of abnormally differentiated CD cells in the absence of renal Ae1. A regulatory feedback of Foxi1 with Ae1 or a multi-protein complex of Ae1 might be required to initiate Pds expression in adjacent cells. The reciprocal signal of a possible Foxi1-Ae1 interaction could induce another unknown factor or again Foxi1, which modulates the expression of Pds in neighbouring cells. A multi-protein Ae1 complex or Ae1 itself might also directly repress the expression of Ae1 in adjacent cells by another unknown signalling cascade, thus providing an autoregulatory mechanism that simultaneously maintains the identity of the effector cell. Similarly, a putative crosstalk of Foxi1 and Cp2l1 might be necessary to repress the expression of CaII in principal cells of the renal MCD, while CaII is strongly expressed in adjacent ICs.

Consistently, the frequency of Cp2l1 expressing cells seemed to increase from the renal cortex towards the inner medulla.

However, with respect to the different signalling cascades that appear to be involved in CD differentiation, the sequential activation of autocrine and paracrine actions seems inevitable. Moreover, this would allow a controlled, rapid and locally organized response to functional demands of the mature kidney.

Embryonic terminal differentiation of the CD cells might nevertheless require additional or different signalling pathways. During embryonic kidney development two populations of precursor cells seem to exist in the CNT and inner medulla, which give rise to types of ICs and PCs. However, the renal CNT originates from the nephrogenic mesenchyme, while the CDs derive from the ureteric bud epithelium. Hence, it is not surprising, that the unusual differentiation of Pds and Aqp2 coexpressing cells was confined to the renal inner medulla. It might well be, that other transcriptional networks and regulators are present in the CNT compared to the renal inner medulla as they derive from different primordia. On the other hand, the urinary composition is constantly modified along the CD system and the composition is different in the CNT and IMCD. Similarly, the interstitial pH is more acidic and the oxygen supply is lower in the inner medulla compared to the renal CNT. Hence, the different functional demands and external stimuli in the renal CNT and inner medulla might influence the interaction of signalling networks and the behaviour of individual molecules.

However, much more work needs to be done to investigate all the different aspects mentioned and to answer all the open questions left.



**Fig. 11.2 Proposed model of renal CD differentiation in mouse.** Notch activity might be modulated by p53 in precursor or terminally differentiated cells. Activated Notch interacts with Cp2l1 and promotes the differentiation of PCs. The transcriptional repressor Cp2l1 inactivates Notch signalling in adjacent cells. Repression of Notch leads to activation of Foxi1, which promotes the differentiation of ICs. Interaction of Foxi1 or another transcriptional regulator with Ae1 or an Ae1 related multi-protein complex induces a feedback loop that drives the differentiation of type B-ICs and maintains the type A-IC phenotype. Cp2l1 might be negatively autoregulated or interacts reciprocally with inactivated Notch, Foxi1, or signalling molecules of differentiated CD cells.

## 12. FUTURE PERSPECTIVES

The present study provides some new and interesting data that suggest a potential role of Ae1 or an Ae1 based multi-protein complex in the terminal differentiation of renal CD cells apart from  $\text{Cl}^-/\text{HCO}_3^-$  exchange. However, further investigations are necessary to establish the speculated role of Ae1 and to characterize the nature of a potential Ae1 related multi-protein complex. In addition, the localization and regulatory role of the transcription factors Foxi1 and Cp2l1 in mouse renal ICs, integrative to data from invertebrate model organisms, raise some new aspects that need further investigation. Respectively, the following experimental approaches are suggestive for future studies:

- Since little is known about potential regulatory pathways and signalling molecules involved in terminal CD cell differentiation, RNA microarray analysis of mature kidneys from Ae1 WT and KO animals could be useful to identify more candidate genes and to further investigate the different signalling pathways (p53, Notch, PI3/Akt/mTOR). Consistently, the mRNA expression of the transcription factors *Gdf15*, *Klf4* and *Bdkrb2*, probably involved in p53 regulated signalling pathway, were already shown to be upregulated in Ae1 deficient mature kidneys.
- Likewise, the protein expression levels, posttranslational modifications, as well as the cellular localization of p53, Gdf15, Klf4 and Bdkrb2 within the kidney of Ae1 WT and KO animals are important to further investigate their role in terminal IC differentiation.
- In the present study, Gdf15 protein abundance was altered in chronic acid loaded mouse kidneys. However, the cellular localization of Gdf15 in the kidney as well as the relevance of the different Gdf15 forms (propeptide, mature and secreted peptide) are not yet known and need to be characterized.



- Good evidence from transgenic mice suggests the Notch signalling pathway critical for CD cell differentiation. Hence, the signalling cascade should be analyzed in Ae1 deficient kidneys with respect to our proposed model (Fig. 11.2). In addition, investigation of Ae1, Foxi1 and Cp2l1 in transgenic Notch mouse models (i.e. Notch deficiency, constitutive activated Notch) would be helpful to further establish their function and proposed interaction in renal IC differentiation.
- A number of Ae1 interacting proteins, critical for cytoskeletal integrity and trafficking events, were already identified in erythrocytes. However, the situation is less clear for the kidney specific Ae1 isoform, since the majority of identified interacting erythroidal proteins do not exist in both tissues simultaneously. Moreover, the kAe1 isoform lacks a portion of the interactive N-terminal domain. Therefore and to proof the hypothesis of an Ae1 based multi-protein complex involved in terminal IC differentiation, a profound analysis of potential interactive partners would be helpful. Technically, several experimental approaches might be applicable such as yeast-two hybrid screening, co-immunoprecipitations or pull down assays with subsequent mass spectrometry analysis.
- Identification of upstream triggers for signalling cascades involved in remodelling of the collecting duct in response to altered electrolyte or acid-base status

- Unfortunately, cultured cells of the renal CD system tend to lose specific protein expression like Ae1, Pds, V-ATPase and dedifferentiate immediately when cultured *in vitro*. Interestingly, the present study revealed the endogenous expression of Foxi1 in untransfected Hek293T cells (Fig. 10.34C). In addition, Bounoure L, Bourgeois S, and Wagner CA noted the presence of the *B1* subunit of *V-ATPase* in these cells (unpublished observation). Therefore, Hek293T cells might be a useful model to study the suggested interaction of Ae1 with putative signalling pathways of terminal cell differentiation. In addition, they could be useful to study a possible interaction of the distinct signalling pathways, i.e. by applying pharmacologic inhibitors of the Notch signalling cascade or siRNA mediated gene silencing. Moreover, two MDCK subtypes were reported with features resembling renal PCs and ICs, respectively [233]. A detailed analysis should investigate their suitability as a model for terminal CD cell differentiation, as well. Alternatively, transfection of cells with transcription factors of interest could provide important information about possible inductive signals (i.e. acid-base, electrolyte composition), posttranslational modifications and activity, interactions with other molecules, their subcellular localization and targeting, as well as suggested interactions with other signalling cascades. Last but not least, the terminal differentiation of renal CD cells might be studied by *in vitro* cultures of mouse embryonic stem cells or metanephric mesenchyme and ureteric bud derived progenitor cells [234, 235].

- The present study provides novel data on Foxi1 and Cp2l1 expression in the mature mouse kidney. Anyhow, these data are preliminary and detailed colocalization studies are necessary to further analyze the expressing cell populations with respect to the distinct CD segments. Furthermore, the cellular distribution and abundance of Foxi1 and Cp2l1 in Ae1 deficient kidneys and acid loaded kidneys needs to be determined. Moreover, immunoblot of Foxi1 on kidney nuclear protein extracts and cytosolic protein fractions suggest multimeric presence of Foxi1 in the distinct cellular compartments. Analysis of the nature and regulatory relevance of these potential Foxi1 manifestations would be necessary to understand the function of Foxi1 in terminal cell differentiation. In addition, a potential interaction of Foxi1 and Cp2l1 needs to be examined and characterized. Consistently, expression of Foxi1 in Cp2l1 deficient kidneys and Cp2l1 expression in Foxi1 deficient kidneys would be useful. Furthermore, the Notch signalling cascade should be analyzed in Foxi1 and Cp2l1 deficient animals.
- In order to definitely exclude dRTA related effects on terminal renal IC differentiation mice deficient for either the B1 subunit of V-ATPase or Rhcg were analyzed in the present study. However, the analysis was performed under standard laboratory conditions where these animals display an incomplete form of dRTA. Therefore, a repetitive investigation of kidneys from transgenic animals receiving a chronic oral acid load *a priori* would give more substantiated results.

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